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**Development of Design Parameters for Virtual Cement and
Concrete Testing**

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Prepared in cooperation with the State of Florida Department of Transportation and the U.S. Department of Transportation.

Approximate Conversions to SI Units (from FHWA)

Symbol	When You Know	Multiply By	To Find	Symbol
Length				
in	inches	25.4	millimeters	mm
ft	feet	0.305	meters	m
yd	yards	0.914	meters	m
mi	miles	1.61	kilometers	km
Area				
in²	square inches	645.2	square millimeters	mm ²
ft²	square feet	0.093	square meters	m ²
yd²	square yard	0.836	square meters	m ²
mi²	square miles	2.59	square kilometers	km ²
Volume				
fl oz	fluid ounces	29.57	milliliters	mL
gal	gallons	3.785	liters	L
ft³	cubic feet	0.028	cubic meters	m ³
yd³	cubic yards	0.765	cubic meters	m ³
NOTE: volumes greater than 1000 L shall be shown in m ³				
Mass				
oz	ounces	28.35	grams	g
lb	pounds	0.454	kilograms	kg
Temperature (exact degrees)				
°F	Fahrenheit	5 (F-32)/9 or (F-32)/1.8	Celsius	°C
Illumination				
fc	foot-candles	10.76	lux	lx
fl	foot-Lamberts	3.426	candela/m ²	cd/m ²
Force and Pressure or Stress				
lbf	pound-force	4.45	newtons	N
lbf/in²	pound-force per square inch	6.89	kilopascals	kPa

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16. Abstract <p>The development, testing, and certification of new concrete mix designs is an expensive and time-consuming aspect of the concrete industry. A software package, named the Virtual Concrete and Cement Testing Laboratory (VCCTL), has been developed by the National Institute of Standards and Technology as a tool to predict the performance of concrete mixes quickly using computer simulation of the hydration behavior of concrete. This software requires thorough characterization of the raw materials of a concrete mix design in order to accurately model the hydration reactions. A two-phase testing program was implemented to evaluate the how well the VCCTL software can predict concrete performance. The techniques required to characterize portland cement were developed and implemented to provide input data for the VCCTL software. The resulting virtual materials were simulated, and a second testing program was performed on physical specimens to evaluate the accuracy of those simulations. The accuracy with which the software simulated basic properties of concrete, such as strength, elastic modulus, and time of set, were examined.</p> <p>The process of acquiring cement phase volume and surface area fraction data has been improved substantially through the use of automated scanning electron microscopy. This has resulted in a more efficient process to obtain cement characterization data for use in the VCCTL software. Comparison of isothermal calorimetry data and corresponding time of set data has shown that a typical Type F high-range water-reducing admixture (superplasticizer) delayed time of set and shifted the main silicate hydration peak by the same amount of time. At the dosages explored within this study, the delay was proportional to the dosage rate.</p> <p>The empirical predictions for compressive strength, which were based on elastic modulus and developed for concretes using coarse aggregates that were mineralogically and/or microstructurally different than typical Florida limestone aggregates, were not accurate for concretes made with Florida limestone. More work is needed to accurately predict compressive strength based on the elastic properties of concrete containing Florida limestone coarse aggregates. A more fundamental approach to the simulation of concrete strength should be investigated. Detailed characterization of the elastic properties of Florida limestones used to produce coarse aggregates for portland cement concrete should be performed. A database of properties of concrete mix designs containing Florida aggregate for use with the VCCTL software and other projects should be created.</p> <p>The VCCTL software was found to be an effective tool for the simulation of elastic modulus of portland cement</p>		

concrete, provided the materials being simulated are properly characterized. The VCCTL software currently does not have a means to incorporate the effects of admixtures on cement hydration. An initial attempt to integrate the effects of a water-reducing admixture, using heat of hydration data, was successful for a Type F water reducer, but the software significantly underestimated the setting time for a Type D water reducer. More work is needed to reliably incorporate the effects of admixtures into the VCCTL software.

There are a number of materials that can be modeled in the VCCTL software that were not considered for this research. There is support for both fly ash and blast furnace slag hydration in the VCCTL software, though the accuracy of the model in this respect is largely unknown. The techniques required to characterize these materials are also more involved due to the significant glassy (amorphous) phase contents of their compositions. The methods by which these materials can be characterized and the accuracy with which they are simulated in the VCCTL software should be explored.

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EXECUTIVE SUMMARY

Background

The development of a tool to predict the properties of portland cement concrete has been the focus of many avenues of research and development. The complexity of the reactions that occur during cement hydration precluded the development of useful computational models of the process until the late 1980s. Over the last 12 years, software known as the Virtual Cement and Concrete Testing Laboratory (VCCTL) has been available for commercial use from the National Institute of Standards and Technology (NIST). This software incorporates microstructural modeling of portland cement hydration, and allows for the prediction of different properties of the hydrated product. The efficacy of the model relies on the proper characterization of the materials being simulated. While the potential usefulness of this tool is substantial, its accuracy, particularly with regards to materials endemic to the state of Florida, has yet to be systematically evaluated.

Research Requirements

Evaluating the accuracy of the VCCTL software requires the comparison of the predicted property values for virtual concrete specimens with the actual property values determined for cured concrete specimens. Elastic modulus and compressive strength for the cured product, and time of set for the plastic product, are the primary predictive outputs for the mix design being modeled. Accurate measurement of these properties for a given concrete is necessary for a valid comparison with the predicted values. Accuracy of the predicted values depends heavily on the accuracy of the input values for the raw materials, which requires that the raw materials are properly characterized. The methods and procedures required for this characterization must be developed and refined.

Research Objectives

The primary objective of this research was to determine the degree of accuracy with which the VCCTL software can predict the various properties of portland cement concrete produced with raw materials typically used in the state of Florida. The specific tasks to meet this objective were as follows:

1. Determine the accuracy with which the VCCTL software predicts the elastic modulus, compressive strength, and setting time of portland cement concrete.
2. Determine if the software can accurately account for the effects of a water-reducing admixture (WRA) on setting time by using isothermal calorimetry data obtained from cementitious samples containing the WRA.
3. Perform sensitivity analyses to evaluate the influence of each material property input on the degree of accuracy with which the above properties are predicted by the software.

A secondary objective of this research was to examine methods of expediting the acquisition of raw material characterization data used for inputs to VCCTL software.

Main Findings

The main findings from this study are summarized as follows:

1. The VCCTL software was found to be an effective tool for the prediction of elastic modulus of portland cement concrete, provided the raw materials used in the simulation were properly characterized.
2. At the current stage of development, the VCCTL software did not accurately predict the compressive strength of portland cement concrete using Florida limestone coarse aggregate. This is expected to be resolved by improving the accuracy of the raw material property inputs, and by making modifications to the software programming.
3. The VCCTL software currently does not have a means to incorporate the effects of admixtures on cement hydration. An initial attempt to integrate the effects of a water-reducing admixture, using heat of hydration data, was successful for a Type F water reducer, but the software significantly underestimated the setting time for a Type D water reducer. More work is needed to reliably incorporate the effects of admixtures into the VCCTL software.
4. The process of acquiring cement phase volume and surface area fraction data has been improved substantially through the use of automated scanning electron microscopy. This has resulted in a more efficient process to obtain cement characterization data for use in the VCCTL software.
5. The empirical predictions for compressive strength, which were based on elastic modulus and developed for concretes using coarse aggregates that were mineralogically and/or microstructurally different from typical Florida limestone, were not accurate for

concretes made with Florida limestone. More work is needed to accurately predict compressive strength based on the elastic properties of concrete containing Florida limestone coarse aggregates.

6. Comparison of isothermal calorimetry data and corresponding time of set data has shown that a typical Type F high-range water-reducing admixture (superplasticizer) delayed time of set and shifted the main silicate hydration peak by the same amount of time. At the dosages explored within this study, the delay was proportional to the dosage rate.
7. Higher dosage rates of Type F high-range water-reducing admixture (superplasticizer) typically resulted in an increase in both elastic modulus and compressive strength, when compared to the control mixes.

Recommendations

Based on the findings from this study, the following recommendations were made:

1. The conclusions drawn from the work performed in this research point to many potential avenues for future research, both to improve the accuracy of the model as well as to explore different material inputs and outputs.
2. The empirical compressive strength model built into the VCCTL software, which is based on elastic modulus, does not accurately predict compressive strength for concrete mixtures which incorporate Florida limestone. Generalized empirical predictions for strength of concrete are inherently limited due to the variability of the materials used in its production. A more fundamental approach to the simulation of concrete strength should be investigated.
3. Detailed characterization of the elastic properties of Florida limestones used to produce coarse aggregates for portland cement concrete should be performed. A database of properties of concrete mix designs containing Florida aggregate for use with the VCCTL software and other projects should be created.
4. The VCCTL software supports the modeling of admixtures if the specific phase surface deactivation behavior of the admixture is known. Methods of obtaining this information, either through material testing, or possibly from the admixture manufacturer, should be investigated.
5. There are a number of materials that can be modeled in the VCCTL software that were not considered for this research. There is support for both fly ash and blast furnace slag

hydration in the VCCTL software, though the accuracy of the model in this respect is largely unknown. The techniques required to characterize these materials are also more involved due to the significant glassy (amorphous) phase contents of their compositions. The methods by which these materials can be characterized and the accuracy with which they are simulated in the VCCTL software should be explored.

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CHAPTER 1. INTRODUCTION

Background

The development of a tool to predict the properties of portland cement concrete has been the focus of many avenues of research and development. The complexity of the reactions that occur during cement hydration precluded the development of useful computational models of the process until the late 1980s due to the limitations of available computational resources. Over the last 12 years, software known as the Virtual Cement and Concrete Testing Laboratory (VCCTL) has been available for commercial use from NIST (The National Institute of Standards and Technology). This software incorporates microstructural modeling of portland cement hydration, and allows for the prediction of different properties of the hydrated product. The efficacy of the model relies on the proper characterization of the materials being simulated. While the potential usefulness of this tool is substantial, its accuracy, particularly with regards to materials endemic to the state of Florida and the simulation of the properties of concrete specifically, has yet to be systematically evaluated.

Model Function

The VCCTL software uses data from real raw materials to create a virtual concrete from which different material properties can be obtained via virtual testing. The model first creates a three-dimensional representation of a portland cement suspension, as would exist at the initial moment of mixing the cement and water. The phase composition of this representation is drawn directly from the data supplied for the cement being modeled. In a process that mimics the actual hydration of portland cement, the model then virtually hydrates this three-dimensional microstructure using a specific set of rules drawn from the observed hydration kinetics and thermodynamics of portland cement. As the virtual microstructure is hydrated, parameters such as heat released, total porosity, degree of hydration, and many others are calculated in real time. When hydration is complete, a finite element calculation, performed on the mesh derived from the virtual microstructure, provides the elastic modulus of the paste. This elastic modulus can then be used in combination with the elastic properties of the coarse and fine aggregate to predict the modulus of the concrete itself. Finally, compressive strength is predicted from this modulus using a simple empirical relationship.

Research Requirements

Evaluating the accuracy of the VCCTL software requires the comparison of the predicted properties of a given concrete with the actual properties of physical specimens. Elastic modulus, compressive strength, and time of set are the easily measured predictive outputs of the model, and knowledge of the actual values of these properties for a given concrete mixture design is essential to ensure accuracy. It is critical that proper characterization of the materials being simulated is performed. This requires implementation and refinement of the existing characterization methods and procedures.

Hypothesis

The Virtual Cement and Concrete Testing Laboratory software predicts the properties of concrete by modeling the hydration of portland cement based on a simulated microstructure. The accuracy with which this simulation takes place is highly dependent upon the accuracy with which the materials being simulated are characterized. The processes by which these materials are characterized may also be made faster and more accurate through the development of new techniques and the utilization of modern instrumentation.

Research Objectives

The primary objective of this research was to determine the degree of accuracy with which the VCCTL software is capable of predicting the various properties of Portland cement concrete. The specific objectives were as follows:

- Determine the accuracy with which the VCCTL software predicts the elastic modulus, compressive strength, and setting time of portland cement concrete.
- Determine the VCCTL software's ability to account for the effects of water-reducing admixtures on time of set through the use of heat of hydration data from isothermal calorimetry.
- Evaluate the influence of different material data on the degree of accuracy with which the above properties are predicted.

A secondary objective of this research was to examine the possibility of expediting the processes required to characterize the different materials being simulated.

Research Approach

The VCCTL software requires detailed information on the input parameters for the material being simulated within the model to provide accurate simulations. The information, termed “inputs” for the purpose of this research, are obtained via the thorough characterization of the materials using laboratory testing methods. Some of this material data is obtained through standard test methods, however the development of specific and complex characterization techniques is necessary for the acquisition of certain material properties. The development and improvement of techniques for the characterization of materials was an initial goal and a significant portion of the work performed for this research. The testing program for the development of material inputs utilized the following material analysis techniques:

- Scanning electron microscopy and energy dispersive x-ray spectroscopy microanalysis ;
- Isothermal conduction calorimetry;
- Laser particle size distribution analysis;
- X-ray powder diffraction (XRD) analysis.

These characterization techniques provided the following inputs for use with the VCCTL software:

- Volume and surface area fractions for the different primary portland cement phases;
- Heat of hydration of portland cement;
- Particle size distribution of portland cement; and
- Mass fractions of sulfate phases in portland cement.

The other major initiative of this research was the investigation of the ability of the VCCTL software to make accurate predictions for different properties of concrete. A study to evaluate the predictions of elastic modulus, compressive strength, and time of set for concrete was constructed to determine the accuracy of the VCCTL software. Secondary aspects of this study included a sensitivity analysis of the software for material inputs, as well as an experimental technique to calibrate the model to include the effects of different admixtures. A physical testing program was implemented to provide information on concrete mixtures against which to compare the results of the model. The following physical tests were performed on fresh and hardened concrete:

- Air Content (ASTM C173);
- Slump (ASTM C143);
- Unit Weight (ASTM C138);
- Temperature (ASTM C1064);
- Time of Set (ASTM C403);
- Compressive Strength (ASTM C39);
- Compressive Elastic Modulus (ASTM C469).

Of these tests, time of set, elastic modulus and compressive strength were compared to predictions made by the VCCTL. Other tests, such as slump, unit weight, temperature, and air content were performed as part of standard mixing procedure for quality control purposes.

CHAPTER 2. LITERATURE REVIEW

Manufacture of Portland Cement

The raw materials required to manufacture portland cement must supply calcium oxide (CaO or C in common cement chemistry notation), silica (SiO₂ or S), alumina (Al₂O₃ or A), and hematite (Fe₂O₃ or F), which are needed to form the four primary phases; Tricalcium silicate (alite, C₃S), dicalcium silicate (belite, C₂S), tricalcium aluminate (aluminate, C₃A), and tetracalcium aluminoferrite (ferrite, C₄AF). The oxides needed to form the four primary phases are typically supplied by some combination of limestone, chalk, slate, and clay, proportioned to give the desired composition on an oxide basis (Mindess & Young, 1981).

Portland cement clinker is manufactured in a rotary kiln, where temperatures at the raw material inlet, at the top of the kiln, are about 450°C, and steadily increase to about 1450° to 1500°C at the clinker discharge at the bottom of the kiln. The mineralogical composition of the cement clinker is determined primarily by the maximum processing temperature, time spent by the raw materials at the maximum temperature, and the rate of cooling. The aluminate and ferrite phases begin to form by solid state diffusion reactions at approximately 1200°C, and the phases then melt at approximately 1350°C. The liquid phase formed acts as a flux, which accelerates the reactions and partially fuses the material into clinker form as it reaches temperatures of at least 1450°C. It is in this latter stage of production where most of the calcium silicates are formed. Portland cement is obtained when the portland cement clinker is interground with gypsum (calcium sulfate dihydrate, CaSO₄·2H₂O, C \bar{S} H₂, where sulfate = SO₃ = \bar{S}), which is added to prevent the concrete experiencing a flash set from the hydration of C₃A (Neville, 2011).

Portland Cement Hydration

When mixed with water, portland cement undergoes a complex set of reactions that result in the transformation of a slurry of cement particles in water to an interconnected solid matrix of hydration products. The rate and resulting products of this reaction are governed largely by the relative concentrations of the four major constituents of portland cement: alite (C₃S), belite (C₂S), aluminate (C₃A), and ferrite (C₄AF). A fifth mineral component, gypsum (C \bar{S} H₂), plays an important role in the early stages of the hydration reaction (Mindess & Young, 1981).

The aluminate and ferrite phases react relatively quickly, but their hydration products add little to the composite strength of the cement. Aluminate is the most readily soluble of the

compounds present in portland cement such that if its proportion is too large, the result will be an immediate stiffening of the cement paste, known as “flash set.” The presence of gypsum within the portland cement is to prevent this flash set by reacting with the aluminate to form insoluble sulfoaluminate (Neville, 2011). The optimum gypsum addition is the amount that will react with almost all of the gypsum, leaving very little aluminate available for direct hydration. In addition to gypsum, other calcium sulfate minerals such as soluble anhydrite (CaSO_4 , $\text{C}\bar{\text{S}}$) or hemihydrate ($\text{CaSO}_4 \cdot 0.5 \text{H}_2\text{O}$, $\text{C}\bar{\text{S}}\text{H}_{0.5}$) can be added to have a similar effect. As with gypsum, the addition of these sulfates must be carefully controlled.

The majority of the strength of hydrated portland cement comes from the hydration of the calcium silicate phases (alite and belite). These phases react with water to produce calcium-silicate-hydrate (C-S-H) and calcium hydroxide [$\text{Ca}(\text{OH})_2$, CH]. While both alite and belite react with water to produce C-S-H, the stoichiometry, solubility and rates of their respective reactions differ. The reaction of alite with water occurs more quickly than that of belite, and is largely responsible for strength development at early ages (up to 28 days). Alite produces a relatively high degree of water-soluble CH relative to C-S-H. Belite reacts more slowly, producing less CH relative to C-S-H, (when compared with alite) and contributing primarily to strength development after 28 days.

Portland cement hydration is a solution-reprecipitation process in which the primary reactants (solid C_3S , solid C_2S , and water) produce secondary reactants (aqueous calcium and silicon ions) that reprecipitate as C-S-H, the binding phase of portland cement. Calcium hydroxide $\{\text{Ca}(\text{OH})_2\}$ is also formed, but it usually precipitates as large crystals in the larger pores and does not have any cementitious properties. The smaller cement particles reduce in size and can be completely consumed because they are more reactive than larger cement particles due to their higher surface-area-to-volume ratios.

The C-S-H tends to precipitate on higher energy surfaces first; that is, areas of high curvature such as large-particle contact surfaces. Thus the larger, slower-reacting cement particles are cemented together and coated by the C-S-H formed by the hydration of the smaller cement particles. As the hydration progresses, there are eventually enough large particles and agglomerates cemented together to form a rigid skeleton that marks the beginning of the setting process. After further hydration, the spaces formed by the dissolution of cement particles and

cement surfaces during hydration, and by the water that is consumed by the cement hydration reactions, are mostly filled with C-S-H and $\text{Ca}(\text{OH})_2$.

Cement Heat of Hydration

The chemical reactions that occur as cement hydrates are exothermic, resulting in the production of heat as the reaction occurs. A typical heat evolution curve is shown in Figure 2-1. The time scale refers to the amount of time that has passed since the mixing of cement and water. The initial spike in Figure 2-1 occurs immediately after water contacts cement, and corresponds to 1) the dissolution of calcium ions and hydroxide ions from the surfaces of C_3S particles which rapidly increases the pH within the system, and 2) the formation of ettringite ($\text{C}_6\text{A}\bar{\text{S}}_3\text{H}_{32}$) from the hydration of C_3A in the presence of gypsum (Mindess and Young 1981), and is followed by a dormant period during which the paste is workable. The dormancy period is typically the time during which placement of concrete would occur. The rate of heat release increases as C_3S and C_2S begin hydrating, indicating the end of the dormancy period. The hydration products link the cement particles together, forming a rigid skeleton, resulting in the set of the paste. The rate of this reaction peaks at approximately 10 hours (Neville, 2011), for normal cement and concrete, after which it tapers off. Approximately 13 hours after the reactions commence, there is a third peak that that occurs which corresponds to a renewed reaction of C_3A following the exhaustion of gypsum (Neville, 2011). The reaction slowly decreases after this third peak, as transport through the crowded microstructure becomes diffusion-controlled, limiting the rate of reaction. The total heat evolved can be found by calculating the area under the power curve shown in Figure 2-1 to obtain a heat of hydration curve, shown in Figure 2-2.

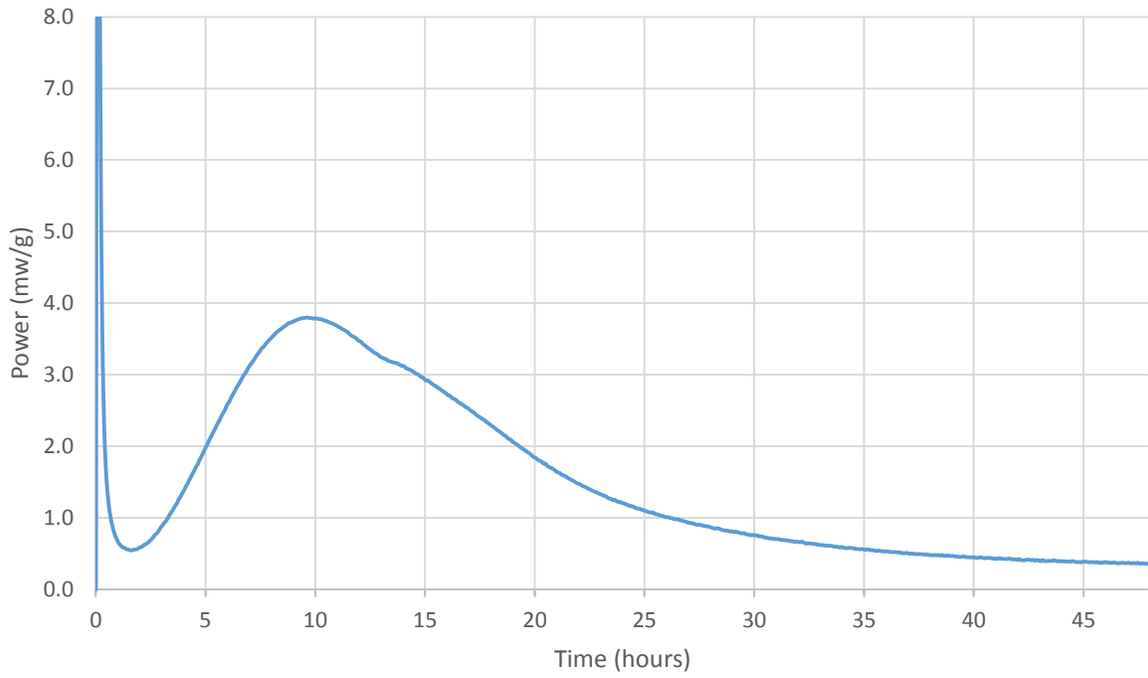


Figure 2-1: Heat Evolution of Portland Cement

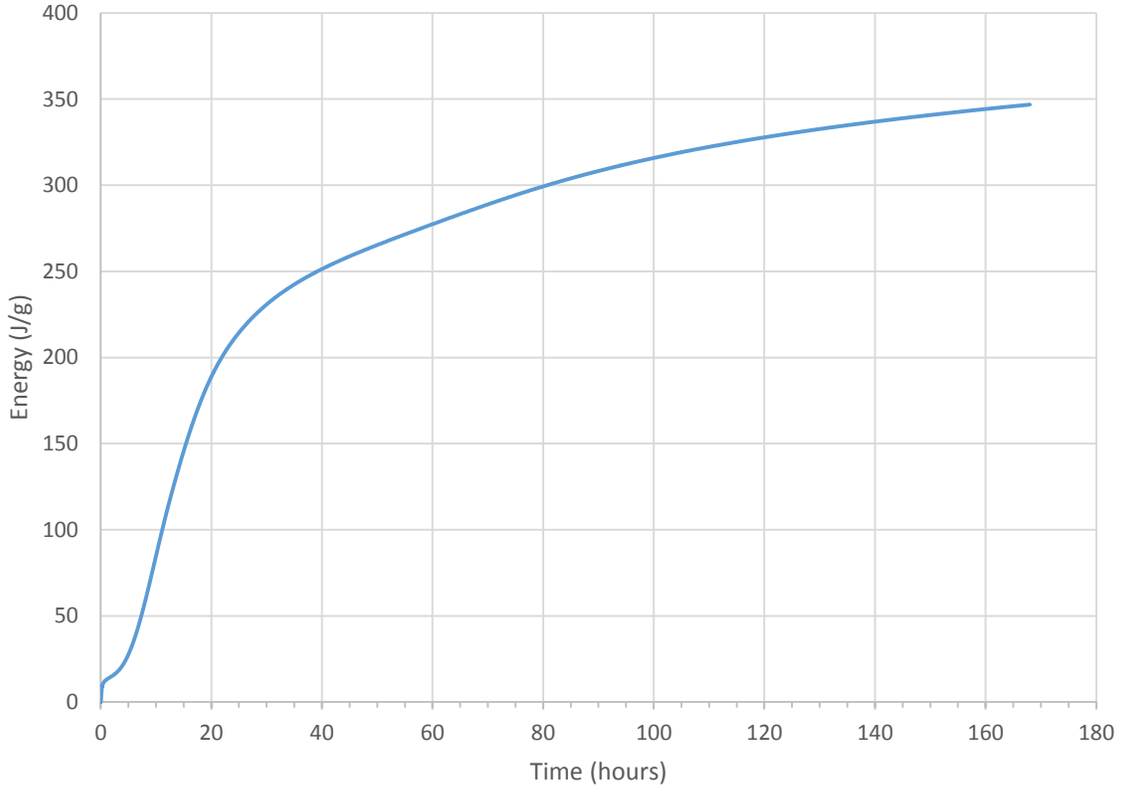


Figure 2-2: Heat of Hydration Curve for Portland Cement

The heat of hydration is a relative measure of the reactivity of the cementitious system or concrete mixture. The chemical composition and fineness of the portland cement govern the rate of heat production and the total heat of hydration of cement and concrete. The standard test methods for the measurement of heat of hydration of cementitious systems are prescribed by ASTM C186 and ASTM C1702.

Admixtures

Chemical admixtures, while not essential for the production of concrete, are used almost ubiquitously to alter the properties of fresh and hardened concrete. The most common types of admixtures are those which improve the wetting characteristics of the solid components of the fresh concrete, which enable the desired plastic properties to be obtained for lower additions of water. The benefit of using a lower water-to-cementitious material ratio (w/cm) is that the hardened properties of the cured concrete are superior due to the formation of a denser microstructure, resulting in a concrete that has a higher compressive strength, lower permeability, and lower residual porosity. These admixtures are referred to as water-reducing admixtures, and operate via a fairly simple mechanism.

Water-reducing admixtures can be divided into two main categories; normal water-reducing admixtures and high-range water-reducing admixtures. The mechanism by which these admixtures operate is fundamentally the same; however their composition and the degree to which their effects are manifested in concrete vary substantially. Normal water-reducing admixtures, classified as Types A or D, typically consist of either lignosulfate or hydrocarboxylic acids. These compounds act by adsorbing onto the cement particles and surrounding them with an envelope of negative charge. This results in mutual repulsion between the particles, causing them to disperse. This manifests on a larger scale as a reduction in viscosity, due to both the mutual repulsion of the particles and the availability of water that would normally be adsorbed onto particle surfaces (Mindess & Young, 1981).

High-range water-reducing admixtures (also referred to as superplasticizers), operate in much the same way as normal water reducers, but to a greater extent. Chemically, high-range water reducers are typically composed of synthesized long-chain organic polymers. These compounds work more efficiently than normal water-reducing agents, exhibiting a stronger negative charge when adsorbed onto the surface of cement particles. The corresponding reduction in viscosity is also more substantial (Edmeades & Hewlett, 1998).

Though normal and high-range water reducers differ in composition and in the degree to which their effects manifest, the similarity of the mechanisms by which they operate results in several common effects. They can have a retarding effect on the initial hydration because the surrounding of the cement particles with admixture compounds temporarily limits the surface area available for reaction to occur. However, due to the more uniform dispersion of the cement particles, more surface area is available for hydration to occur, which can result in higher strengths at early ages after the temporary retardation (Neville, 2011). Normal water reducers that exhibit this temporary retardation are classified as Type D, while those that do not are classified as Type A.

Strength of Concrete

The most frequently used industry metric for the evaluation of a concrete mix design is strength. Strength of concrete is most commonly taken to mean uniaxial compressive strength, which is typically obtained via the crushing of cast cylinders. The strength of the concrete is influenced by a number of factors, including the relative concentrations of the different phases of portland cement; however, the water-to-cement ratio (w/c) is typically regarded to have the greatest effect on strength of a portland cement system due to two primary factors. The more important of these is a reduction of the gel-to-space ratio ($gel/space$), which can be described as the ratio of the volume of hydrated cement paste to the sum of the volumes of the hydrated cement and capillary pores. This can more simply be thought of as the density of the cement paste, as it follows that the more lower-density water that is present relative to the much denser cement, the lighter the resulting paste will be. $gel/space$ and w/c are inversely related, that is the lower the w/c , the higher the $gel/space$. As the cement hydrates, water in the capillaries is consumed, leaving porosity. As w/c decreases and $gel/space$ increases, porosity in the cured concrete decreases, resulting in an increase in strength.

The other main mechanism by which w/c influences the strength of concrete is related to the relationship between the coarse aggregate and the hydrated cement paste. As w/c increases, bleeding (separation of the water from the paste) begins to occur around the aggregate particles. This bleeding results in cracks around the aggregate particles that lower the required stress to cause failure (Maso, 1996). The extent to which this phenomenon occurs depends on the amount of water in the paste, with lower water contents being less affected.

Scanning Electron Microscopy

Scanning electron microscopy (SEM) is a microanalysis technique that allows for the imaging of features several orders of magnitude smaller than those that can be imaged using conventional optical microscopy. The fundamental operation of a SEM relies on the interaction of a focused electron beam with the surface of the specimen being imaged. This electron beam is typically created using a thin tungsten filament, which when subjected to heat and a very strong accelerating voltage, results in the emission of electrons. The electrons are focused using a series of lenses and apertures to a spot on the surface of the specimen. A schematic of a typical instrument is shown in Figure 2-3.

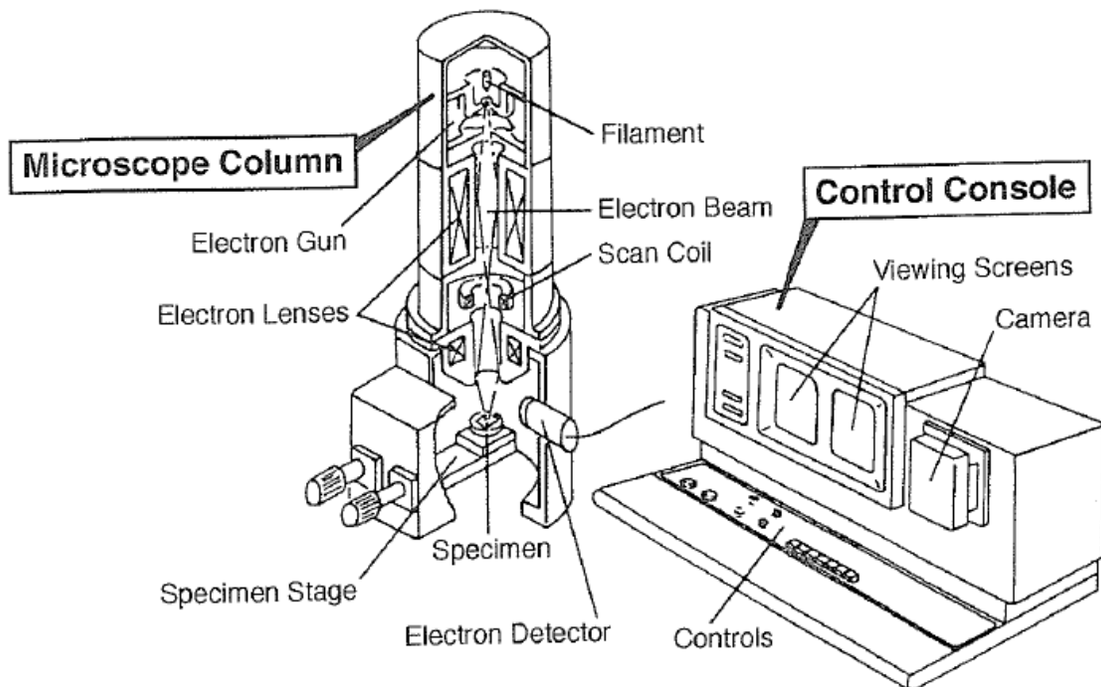


Figure 2-3: Scanning Electron Microscope (Goldstein, et al., 2007)

The focused electrons penetrate into the surface of the sample and collide with the atoms present. The region in which this occurs is referred to as the interaction volume, a diagram of which is shown in Figure 2-4.

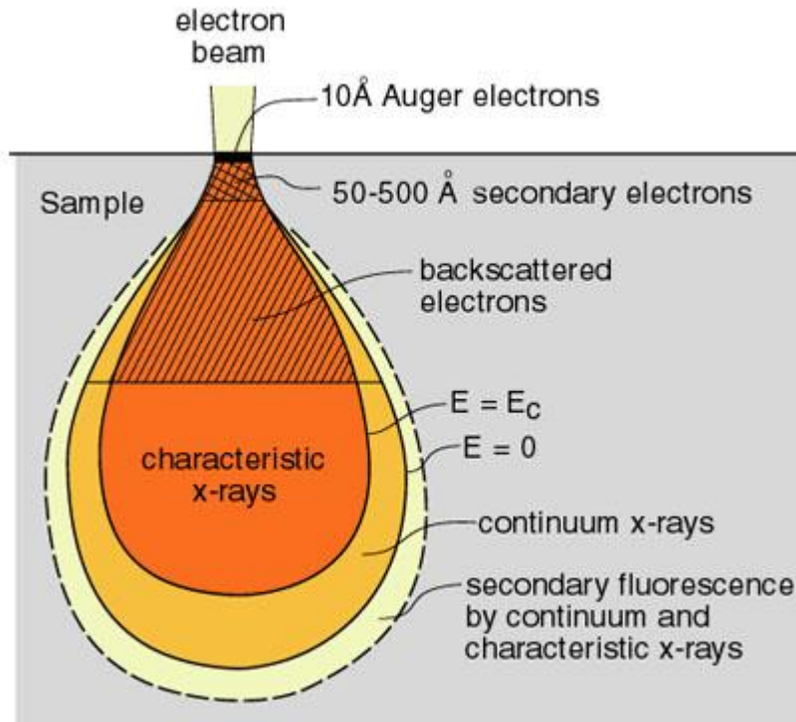


Figure 2-4: Beam Interaction Volume Showing Different Signal Emission Regions (Wittke, 2008)

The interaction of the electrons with the atoms in the target specimen result in several different types of emissions that can be collected and used to create an image, of which, two primary types of emitted electrons are used for image creation. Backscattered electrons (BSE) are emitted from the sample when electrons from the beam undergo elastic collisions with the atoms in the sample and are scattered back out of the surface of the sample. These electrons have high kinetic energy and are detected by an annular detector that surrounds the aperture from which the beam originates. The other type of electrons that are emitted from the sample are secondary electrons (SE), which undergo inelastic scattering events with the atoms in the sample and lose much of their kinetic energy in the process. The electrons that have sufficient energy to make it back to the surface of the sample are attracted toward a positively-charged detector.

Characteristic x-rays are emitted from the interaction volume as a result of inelastic scattering events. These photons have characteristic levels of energy depending upon the element

with which they interact and enable the elemental analysis of a specimen, and this form of imaging is referred to as energy dispersive x-ray spectroscopy or EDS.

The image in a SEM is produced by collecting the signal of interest from the target area, and moving the electron beam rapidly in a grid pattern. Different signal intensities are created at each spot (corresponding to a pixel) to produce contrast within the image. Contrast mechanisms of the different imaging modes are the result of the physiochemical properties of the specimen. BSEs are produced as a result of elastic scattering events, which occur more frequently in specimens containing elements with relatively high atomic numbers. This results in an image in which the brightest areas depict regions that have the highest average atomic number. Topographic irregularities on the surface of the sample, which have exposed edges that provide relatively short distances for secondary electrons to escape, appear brighter due to the larger numbers of escaped secondary electrons. This “edge” effect enables relief or scratches in the sample surface to be easily seen.

Though SEM allows for higher magnifications and different imaging modes than conventional optical microscopy, certain aspects of the imaging process limit the types of samples that can be imaged. The whole SEM is under vacuum, as gas would ionize in the electron beam if present. This precludes the imaging of samples containing water or any other compounds that would boil away at very low pressures. The sample must also either be conductive or be made conductive, otherwise electrons will charge the surface of the specimen, resulting in bright artifacts in the image. For elemental analysis using EDS, the sample must be reasonably flat, as topographic variations can influence the degree to which x-rays are emitted from different parts of the sample.

Computer Modeling

Since the invention of the transistor and the subsequent exponential growth of computational processing power, efforts have been made to simulate complex systems through the use of computer models. A model can be quite simply defined as a representation of a system upon which operations can be performed to examine the response of the system to specific stimuli. The engineering applications of computer modeling typically extend to numerical or stochastic simulations, the former describing a system governed by equations that cannot be solved analytically, and the latter describing a system where the occurrence of events is probabilistic in nature. Both types of simulations have applications in civil engineering, with

numerical modeling of structural systems using the finite element method being most common, while stochastic simulations are frequently applied to the simulation of weather phenomena.

The finite element method is a modeling technique used to numerically simulate the overall behavior of geometrically complex objects by discretizing them into “elements” that are interconnected, each of which can be described with its own set of equilibrium equations. Given a certain set of “boundary conditions” which act on the geometry, the system of equations from the individual elements can be solved to determine the influence of the boundary conditions on the entire system (Zienkiewicz, et al., 2005). A common example of the application of this method is the analysis of the deflection of a truss structure with pinned connections subjected to specific loads. In the event the modeled structure remains in static equilibrium, the forces in each discretized element of the structure must also be in equilibrium. Since the relationship between force and displacement for each element is known, the system of equations for all elements can be solved to find the resulting displacement of the structure as a whole.

Stochastic modeling is used to simulate the behavior of systems that are non-deterministic, which means that they exhibit behavior that contains an element of randomness. These systems can instead be described by the probabilities that events within the system will occur. For example, the range of particle sizes that make up portland cement is highly variable. Given a finite number of particles for a particular cement, the probability that a particle will be of a particular size (the event) can be obtained by dividing the number of times that a particle, that falls within a given size range, is observed, by the number of particles that are measured.

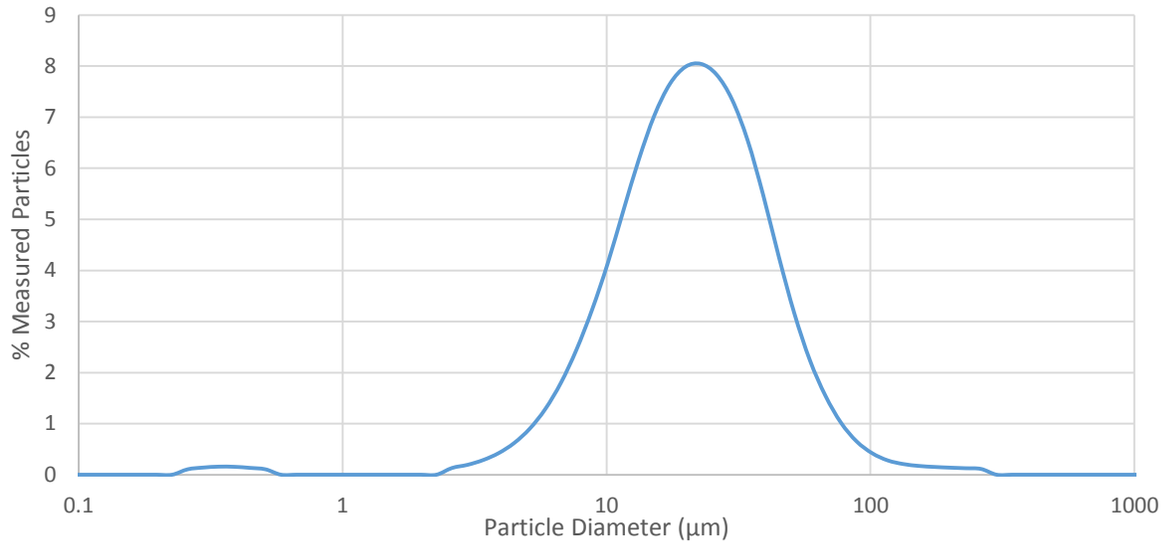


Figure 2-5: Particle Size Distribution

If all particles in a cement sample are measured and classified into different size ranges, the frequency of occurrence of individual particles can be plotted for each size range creating the distribution presented in Figure 2-5. The result is described as the probability density function for the sizes of portland cement particles. This information can now be used to create a set of virtual cement particles with the same distribution of sizes.

Cementitious Simulation

Background

The fundamental goal of modeling cementitious systems is to predict the physical and durability properties of the material. Important structural-related properties of the hardened cementitious microstructure modeled include hardened properties such as modulus of elasticity and compressive strength, as well as durability-related properties such as porosity and permeability. The creation of long-lasting concrete structures relies on the knowledge of these characteristics of the material. The accurate determination of these properties via computer simulation is dependent on the creation of a realistic virtual microstructure from which they can be measured. Since the properties of the microstructure are dependent upon the composition and hydration conditions of the cement from which it is created, there must also be a method by which the development of the microstructure over time can be simulated.

History

The origins of cement microstructural modeling are from research performed in the 1970's on the structure of amorphous semiconductors (Garboczi et al., 2000). Initial attempts to calculate structural properties using approximate analytical solutions were only marginally successful until a computer model was constructed consisting of several hundred randomly linked atoms. Operations were performed on this model to calculate properties, and the resulting properties were compared to experimental results. This model represented one of the first attempts to model amorphous materials computationally at the atomic level.

The first attempt to model concrete computationally was recorded in the publication by Wittmann, Roelfstra, and Sadouki in 1984 on the numerical simulation of the structure and properties of concrete in two dimensions (Wittmann et al., 1984-1985). The two-dimensional silhouettes of aggregates were first characterized by transforming the contour of an aggregate particle section into polar coordinates. The radius of the particle as a function of the angle θ about the y-axis could then be plotted, and the resulting frequency distribution obtained. The frequency distributions of several particles of a given aggregate normalized for size were combined to form a "morphological law" which was then used to generate aggregate sections computationally.

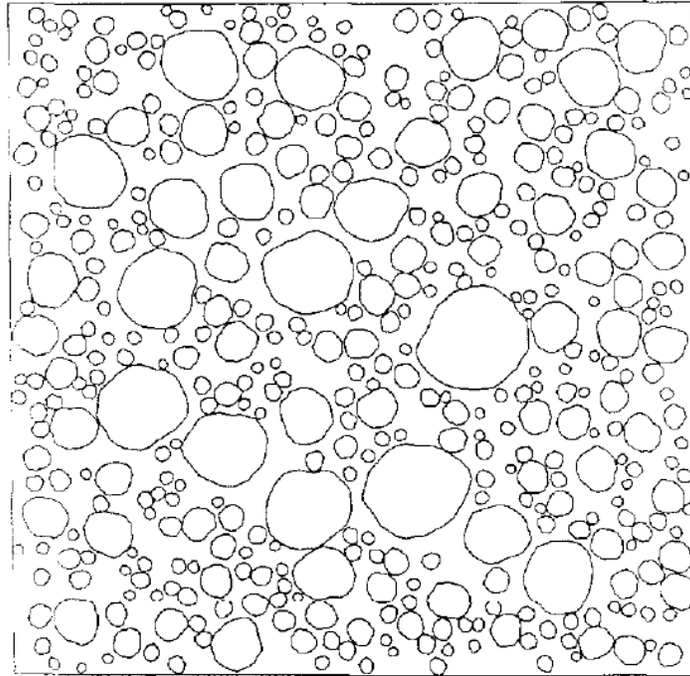


Figure 2-6: Computer Generated Concrete Structure Using “morphological law” (Wittmann et al., 1984-1985)

The computer generation of the two-dimensional section of a composite concrete matrix was created using the morphological law in combination with a measured aggregate size distribution (Figure 2-6). The research addressed the possibility of generating a three-dimensional concrete structure using the same principles, but ruled out the analysis of a three-dimensional structure due to the computational limitations of typical computers during that time. An alternative approach was used to represent the three-dimensional composite structure using two-dimensional images. To obtain a two-dimensional image representative of the three-dimensional structure, the size distributions of aggregate particles for a number of arbitrarily chosen planes from the three-dimensional structure were averaged and the resultant distribution was then used to create a representative two-dimensional composite structure.

The two-dimensional structure was represented by a finite element mesh, which combined with an assumed value for the modulus of elasticity of the cement paste, allowed for the numerical computation of the elastic modulus of the composite as a function of aggregate modulus of elasticity. Different meshing techniques allowed for the computation of the effective diffusion coefficient of the composite structure. Finally, the research by Wittmann et al.

concluded that further work could be done in which a mesh containing more detailed material properties could be used to predict more complex behavior, such as creep, shrinkage, and non-linear stress-strain behavior.

In 1986 the details of a three dimensional hydration model for C_3S were published (Jennings & Johnson, 1986). A model was created which followed rules based on measurable characteristics of the system being simulated. The primary focus of this model was not to accurately simulate the hydration behavior of C_3S , but instead to provide a tool that connected the probable mechanisms by which the reaction occurred with the measurable behavior of the system. Such a tool is useful for the evaluation of the validity of a proposed mechanism by simulating its effect on the behavior of the system.

The model simulated C_3S particles as spheres, with the size distribution, number, and initial packing type of the spheres entered by the user. Other user-controlled inputs included the density of different hydration products, rules governing the distribution of hydration products, and the rate-controlling reaction step at each stage. Simulations within the model began by randomly distributing the specified number of particles with the specified size distribution. Simulation of the cement hydration was initiated at the location of the smallest particle and sequentially incremented. The diameter of the smallest particle was reduced by an amount that was based on information known about the specified rate-controlling step. The space left by the reduction in diameter was filled with hydration product surrounding the particle, and any remaining hydration product was added to the diameter of the particle. This process was repeated for each particle in sequence. One hydration cycle was complete when all particles had undergone this process. The simulation continued until either all anhydrous phases were consumed, the thickness of the hydrated product layer reached a user-specified value, or a specific number of cycles were completed.

The resulting microstructure derived from simulations with this model was compared to SEM micrographs of the hydrated C_3S grains. A promising degree of resemblance was observed. The future objectives of this research envisioned the use of the model for the evaluation of different mechanical properties of the composite structure, and predicted the model would enable research into proposed reaction mechanisms. The incorporation of the other phases of portland cement was also envisioned.

This model was used as a tool for the simulation of cement hydration products for several years, including work to apply a random walk algorithm for continuum models (including the Jennings and Johnson model) in order to compute the electrical and diffusive transport properties of the microstructure (Garboczi et al., 2000). This work introduced the idea of digitizing the microstructure and experimented with a number of computations on a continuum-based microstructure model. It was during this time that the advancements in computer technology made it possible to create model simulations of sufficient size. The concept of a digital microstructure is essentially a three-dimensional (3-D) image consisting of voxels (3-D pixels). Each voxel represents a discrete volume of material with specific properties. A virtual microstructure is created out of the voxels based on the physical and chemical characteristics of the cement being modeled. Once the input parameters have been entered into the computer for simulation, the virtual microstructure undergoes the simulated hydration process, with each voxel acting as an independent agent. Voxels follow specific rules for dissolution, diffusion, and reaction based on their phase and the known thermodynamic and kinetic behavior of cement hydration. The creation of this digital microstructure led to the realization that nearly any finite element or finite difference algorithm can be applied to measure the properties of the microstructure. The only remaining steps were to establish the applications of percolation theory and composite material theory to the digital microstructure.

The interconnected porosity in a cementitious body can be considered to be a system of large pores, formed by the open interstices of particles in contact, that are connected by tortuous, sheet- or capillary-like continuous spaces formed by close particle contact areas. The rate of diffusion or fluid transport of material through the cementitious body is related to the volume of the porosity, the tortuosity of the porosity, and the concentrations and concentration gradients of the materials in the fluid. Percolation generally refers to the movement of a fluid through small but interconnected porosity. However, when applied to the numerical modeling of a cementitious matrix, percolation theory is based on the connectivity of random phases in a multiphase environment (Garboczi & Bentz, 1998).

There are two primary applications of the concept of percolation to the measurable properties of a hydrating microstructure. The first is to model the degree of hydration of the microstructure which is dependent on the extent with which reactants and reaction products can diffuse throughout the microstructure through the water-filled capillary network. The ease of

transport is dependent on the size (equivalent diameter) and degree of interconnectedness (degree of percolation) of the capillary network. As hydration progresses, the reactants and products diffuse through the water-filled capillaries and deposit on the surfaces of the interconnected capillary pathways, gradually reducing their size. The reduction in capillary size creates an increase in tortuosity and a cementitious system with a lower rate of diffusion which ultimately results in a reduced rate of transport. Thus, the degree of percolation of pore space at early ages affects the extent and rate of initial hydration, and the progressive reduction in the rate of diffusion reduces the progressive rate of hydration as the capillaries constrict.

The application of percolation theory to a digital lattice was established through work on the conductivity of a plane containing random holes (Garboczi et al., 1991), and an algorithm was developed to compute the linear elastic properties of a heterogeneous material from a digital lattice a few years later (Garboczi & Day, 1995). The combination of different digital lattice-based methods for the measurement of cement microstructure led to the eventual development in 1997 of a three-dimensional cement hydration model known as CEMHYD3D, the model, which in an updated form, underpins the VCCTL software.

The VCCTL software was introduced by the National Institute for Standards and Technology (NIST) in 2001 and was intended as a unified solution for modeling concrete at multiple length scales. Concrete can be described as a multi-scale material, in that the properties of the paste microstructure, which exist at a micro (10^{-6}) scale, influence the properties of the paste at a millimeter scale, which in turn influences the properties of the concrete at a macro-scale level. The intention of the VCCTL software was to be a start-to-finish modeling solution for concretes and mortars, requiring only data on the materials being simulated to provide accurate measurements of mechanical and transport properties.

Validation of CEMHYD3D

CEMHYD3D, the hydration model used by the VCCTL software, has been available for public use since 1997. There are a number of publications of note from this period, some of which include a quantitative comparison of the initial and hydrated microstructures generated by CEMHYD3D (Bentz 2005), and an investigation of the influence of ground limestone filler on cement hydration (Bentz 2006). The research performed by Bentz, compared the simulated with actual cement microstructures in an effort to investigate the validity of CEMHY3D for the modeling of hydrating portland cement microstructures. Microstructures were compared visually

and also quantitatively using a normalized two-point correlation function. The initial microstructure of a cement was compared to model simulations using spherical particles as well as real particle shapes. The visual comparison of the different initial microstructures can be seen in Figure 2-7.

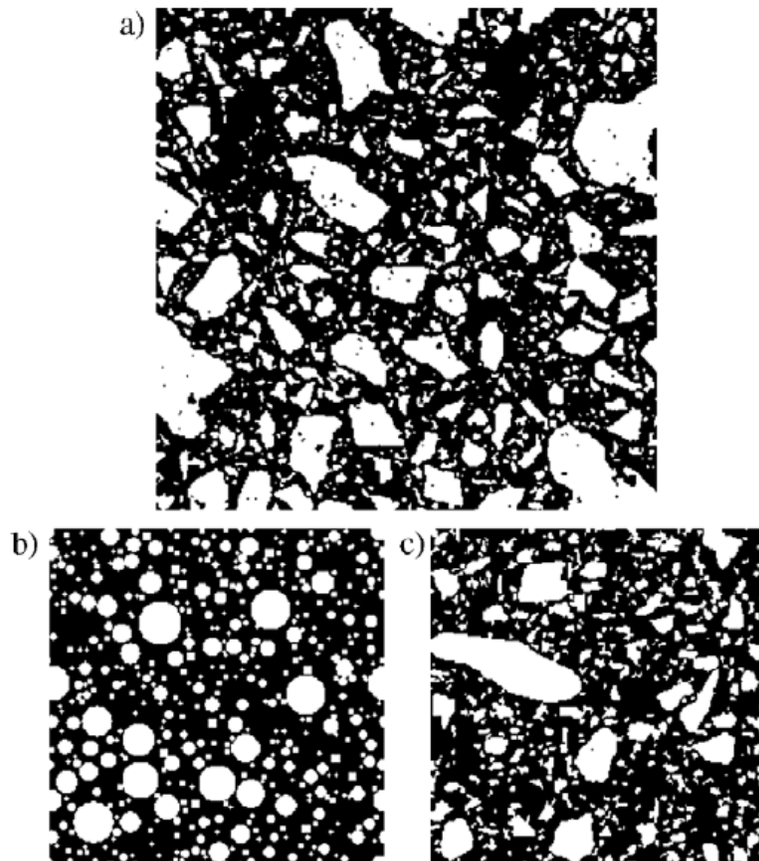


Figure 2-7: Initial Microstructures of (a) Real Cement, (b) Model with Spherical Particles and (c) Model with Real Particle Shapes (Bentz 2006)

The images in Figure 2-7 were compared mathematically to evaluate the spatial accuracy of the simulations using a two point correlation function. Similar comparisons were made between real and simulated hydrated microstructures. Conclusions made from this research were that CEMHYD3D agreed very closely with the real systems, both in the visual comparison of model and real microstructures, and the comparison of two-point correlation functions for model and real microstructures (Bentz, 2006).. It was also observed that the model microstructures with

real particle shapes agree better with the two-point correlation functions of the real cement, and that larger model system sizes also improve the agreement between these functions.

The investigation of the influence of finely ground limestone on the hydration of cement (Bentz D. P., 2006) was triggered by a change in the ASTM C150 standard specification for portland cement to allow for the inclusion of up to 5% of ground limestone, and the claim that the presence of limestone does not influence the overall performance of portland cement. CEMHYD3D was modified for this study to account for the hydration behavior of cements containing ground limestone. The main modifications to the model accounted for the potential for the additional surface area contributed by the limestone to provide more nucleation sites for the growth of hydration products, as well as the potential for a chemical reaction to occur between the cement and limestone itself. The results of the hydration model were validated against experimental measurements of degree of hydration via non-evaporable water content. This study concluded, based on good agreement of model and experimental data, that CEMHYD3D successfully incorporated the chemical and surface area effects of ground limestone.

Existing Research Using the VCCTL Software

The VCCTL software has been available for commercial and research use for over a decade, and in that time has been continually developed and improved. Because of the continual improvements made to the model during this time period, preference was given to more recent research into the applications of the software. One of the simplest applications of the VCCTL software is as an exploratory tool to investigate the effects of changes to cement chemistry. The ability of the model to consider changes not only to cement composition but also fineness (via particle size distributions) allows for optimization of the cement being produced with a minimum of additional labor.

Work done by Siam Cement Group (SGC) (Sahachaiyunta, 2012) investigated the influence of sulfate content and fineness on mortar strength. Twenty-five combinations of fineness and sulfate content were simulated at 3, 7 and 28-day ages. The strength of the mortars was calculated for all twenty-five combinations at different ages and used to evaluate the performance of each permutation. It was concluded from results of plant trials that the VCCTL software could be an effective tool for the optimization of different cement production

parameters, especially when considering more complex blended cements or the implementation of high risk (economically) production strategies.

The VCCTL has great potential as an optimization tool for production within a cement plant. Mapei Industries in conjunction with the University of Padua have performed some successful validation studies using the VCCTL (Valentini, et al., 2013). The accuracy of both the hydration simulation and modulus calculation of the software was evaluated for two different cements. The experimental program consisted of the following:

- In situ x-ray diffraction was used to measure the difference of volume fractions of different cement phases and resultant hydration products over time.
- Elastic modulus of mortar cubes was obtained and elastic modulus, compared to the predictions / calculations by the VCCTL with respect to time.
- Additionally, a power law equation was developed to empirically predict the strength of mortars from elastic modulus.

Valentini et al. determined that the VCCTL software simulates the hydration behavior of the chief hydration products, C-S-H and CH, and the diminishing quantity of C_3S with a reasonable degree of accuracy. However, there were substantial differences between the measured and predicted volume fractions of ettringite within the system. This was attributed to poor simulation by the VCCTL of the increased thermodynamic stability of ettringite in the presence of carbonates. It was concluded that the inaccuracy would be largely irrelevant for the prediction of mechanical properties, as ettringite made up less than 3% of the total volume of the cementitious system. The study determined that there is relatively strong agreement between prediction and measurement of elastic modulus and the empirical relationship developed for the prediction of strength from modulus also correlated well with measurements. Of note is the strong caution against the use of the empirical equation for strength developed in this study for mortars in general, as the properties of the input materials may vary substantially.

Another current focus of research being conducted using the VCCTL software is the characterization of the influence of addition of liquid admixtures. The software has the capacity to integrate the influence of admixture additions into the system provided the behavior of the admixture is known with respect to cement phase surface deactivation time, which is a parameter that is not readily available for the majority of admixtures (Bullard & Obla, Virtual Testing of Ready-Mixed Concrete, 2004). Previous research has examined the behavior of admixtures,

using gel permeation chromatography. Analysis of liquids obtained from filtering cement and pure cement phase pastes enabled the measurement of the quantity of superplasticizer adsorbed by the cement or constituent cement phase. The total quantity of plasticizer can be determined based on the specific surface area of the cement particles obtained from Brunauer-Emmett-Teller (BET) gas adsorption. This then allows for the determination of the duration of deactivation of cement particle surfaces, which can then be entered directly into the VCCTL software (Russo, 2012). This technique is still in development, and has yet to be verified.

There is no research currently published that validates the accuracy of the VCCTL software to predict the behavior of concrete as opposed to mortar. Studies do support the accuracy of the software in the prediction of the properties of the cement paste as well as mortars made with that paste (Bentz, 2006) (Valentini, et al., 2013) (Sahachaiyunta, 2012); however, the methods by which the software incorporates coarse aggregate have yet to be systematically validated.

Current Limitations of the VCCTL Software

Though the VCCTL software is the product of years of research into the modeling of cementitious systems, there are still some aspects for which it is not robust. There is little research which involves the simulation cementitious systems which include large percentages of supplementary cementitious materials (SCMs) such as fly ash or ground granulated blast furnace slag (slag). Fly ash and slag are products that are used in almost all concrete mixture designs today. The Florida Department of Transportation, for instance, has a minimum fly ash requirement for all structural concrete mixture designs. The software has some ability to handle these materials; however, it has not been very well validated and is missing a number of details as far as the thermodynamics and specifics of reactions are concerned (Garboczi, et al., 2010). The VCCTL software has limited support for admixtures, and it is implemented in a way that makes them difficult to implement into the simulation of cementitious systems. The model requires that the specific amount of time that the surface of a phase is deactivated by the admixture is known. This information is difficult to obtain and may only account for certain aspects of admixture behavior. Efforts are currently underway to develop a process by which admixtures can be characterized to obtain this information. It may also be possible to obtain this information from the manufacturers of the admixtures themselves.

The long-term durability is one of the most important aspects of concrete, both for the purposes of sustainability and economy and thus, a model that can predict the long-term performance of concrete would benefit from some sort of durability prediction and degradation modeling. Currently, the VCCTL software does not provide direct functionality for the prediction of durability in concrete. The hydration model provides several outputs that can be used for additional modeling to simulate the durability of concrete, however no direct outputs currently exist.

Other Cementitious Modeling Software

The prediction of concrete performance is a problem for which a solution has great economic potential. As a result, several commercial modeling solutions for concrete exist. None of the models provide results that are based on the fundamental processes underpinning cementitious systems, such as those simulated by the VCCTL software, but their more specific focus has enabled viable alternatives.

The software model STADIUM is modeling software for cementitious systems that is used to make service life predictions for concrete mixtures based on material testing. Thus, a concrete mixture design being considered for use can be examined via physical testing and compared to large databases of similar mix designs evaluated under different exposure conditions. The model has been used for the prediction of service life for many different mixture designs in many different environmental conditions, and has seen use by organizations such as the US Navy. Though the service life predictions of STADIUM have reached a degree of commercial acceptance, the models and databases from which it makes these predictions are proprietary. The exact details of the function of STADIUM are not documented.

Mines ParisTech has produced a numerical model to simulate the effects of cracking and alkali silicate reaction (ASR) in cementitious systems (Peyrot, 2006). The model uses a finite element approach to simulate the growth of reactive aggregates bound in a cementitious matrix, and the resulting bulk swelling. This model incorporates methods for simulating crack development and propagation in heterogeneous materials. Though this direction of research is promising, this model is still in the research stages, and has not seen any commercial use.

Another established concrete material model that has a somewhat different application is TNO DIANA, a finite element model for reinforced concrete structures using realistic material properties for concrete and steel. The software is capable of simulating cracking and crushing

failures, concrete-reinforcement interface zones, creep and shrinkage-induced cracking, material aging effects, and thermally induced stresses. Work performed at the University of Florida expanded the capability of the software to modeling of temperature rise in mass concrete structures due to heat generated during cement hydration (Tia, et al., 2013). The use of heat of hydration data obtained via isothermal calorimetry to build input files for TNO Diana enabled the prediction of thermal stress distributions and cracking in mass concrete structures. Comparison of the results of the simulations with real mass concrete structures found that the prediction capability of the model was fair, with the model predicting higher temperatures than those encountered in the field.

Currently, the Massachusetts Institute of Technology (MIT) Concrete Sustainability Hub is working in cooperation with the Portland Cement Association (PCA) to create and model the atomic structure of portland cement concrete. The goals of this Concrete Sustainability Hub are to bring about breakthroughs in concrete science, and transfer those advancements to the concrete industry (Who We Are, 2012). One of their key goals is the reduction of CO₂ emissions from portland cement production. Currently a number of advances in the understanding of the nano-scale atomic structure of certain phases in cement have been realized, but a practical link between any of these advances and the modeling of portland cement-based systems has not been documented to date.

The specialized nature of these models differentiates them from the VCCTL software, which is intended to be a full spectrum solution for the modeling of concrete. The further development of the software to incorporate features such as fracture analysis and service life prediction is ongoing.

CHAPTER 3 VCCTL SOFTWARE MATERIAL INPUTS

Overview

The most important part of using the VCCTL software is the proper characterization of the materials being simulated. The model takes a number of different types of input data, some of which are essential to running a simulation, and others which simply serve to enhance the accuracy of simulations, or alter the way in which the virtual microstructure is generated or hydrated. The inputs that are required for any hydration simulation include the following:

- Cement phase volume and surface area fractions;
- Cement phase relative spatial distributions;
- Cement particle size distribution.

The following inputs are required to perform simulations including aggregates (mortar or concrete),

- Aggregate bulk and shear moduli;
- Gradation of aggregate;
- Specific gravity of aggregate.

There are other inputs that are not required by the VCCTL software to run simulations; however, it is possible that the presence of these inputs may improve accuracy. For the purposes of this research these inputs have been deemed “non-essential” and are as follows:

- Heat of hydration curve of portland cement;
- Mass fraction data of calcium sulfate phases present in cement;
- Particle size distribution of sulfate phases;
- Shape data of aggregate;
- Shape data of cement particles;
- Simulated microstructure size;
- Particle dispersion of cement within microstructure.

The VCCTL software will accept and include data on the composition and particle size distributions of pozzolanic materials such as fly ash or slag, as well as the chemical surface deactivation behavior of admixtures. However, the investigation of these properties was out of the scope of this research project.

Model Inputs for Portland Cement

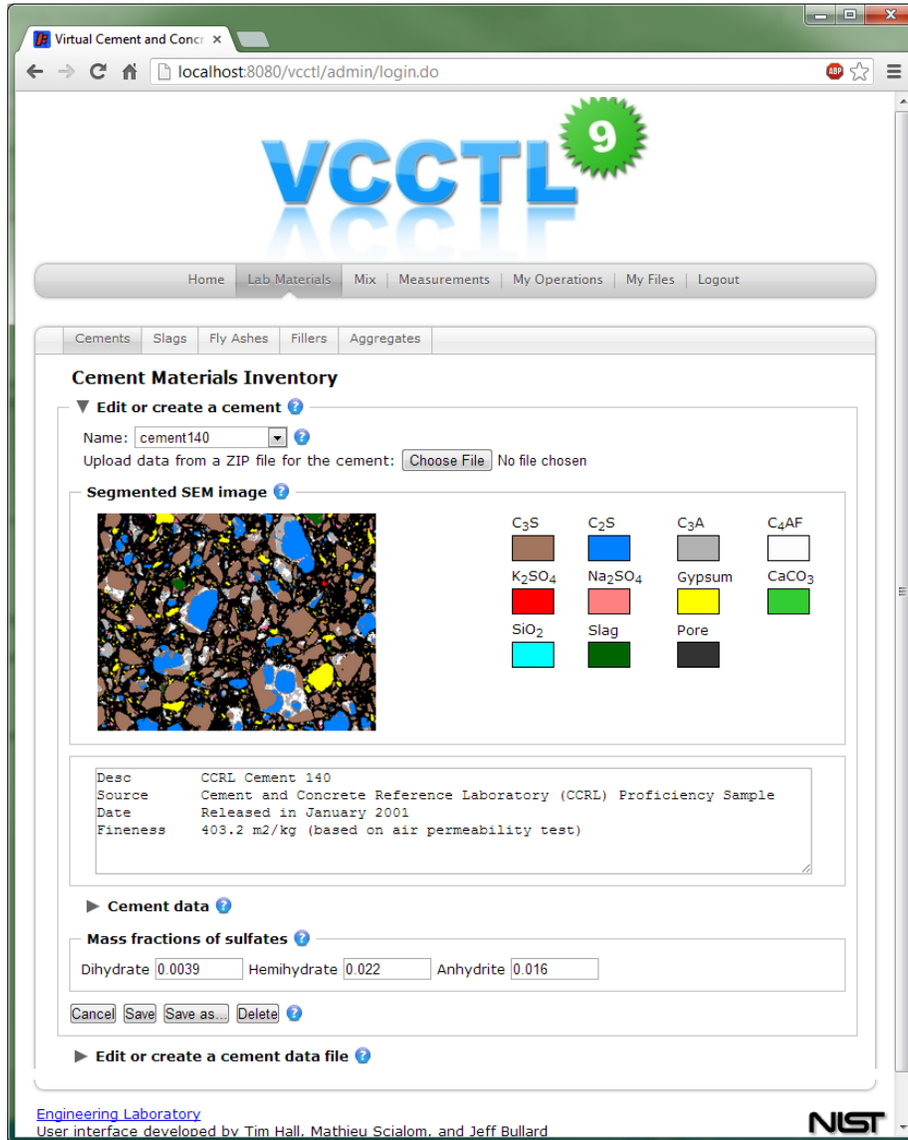


Figure 3-1: Cement Input Screen of the VCCTL Software

The inputs required for portland cement can be divided into two categories: inputs that are used to create the digital microstructure, and inputs used for the virtual hydration of the cement microstructure. Many of these inputs can be obtained using the procedures as prescribed in standardized testing methods and/or with specialized equipment, while others require more complex analysis techniques and procedures. The cementitious material inputs are entered into the VCCTL software through the Lab Materials – Cements page, shown in Figure 3-1.

Microstructural Inputs

The following inputs are used by the VCCTL software to create a digital microstructure that is representative of the cement from which they are obtained.

Gypsum Phase Mass Fractions

ASTM C1365 is used to determine the mass fractions of the different crystalline phases present in portland cement. VCCTL software uses the data produced by this test to obtain the mass fractions of the different calcium sulfate phases, the input fields for which are visible in Figure 3-1. Calcium sulfate or gypsum is added to cement clinker during grinding to prevent the flash set of C_3A during hydration. There are three possible manifestations of gypsum, which are distinguished by the molar ratio of water to calcium sulfate, with hemihydrate having a ratio of 0.5 to 1, dihydrate having a ratio of 2 to 1, and anhydrite having a ratio of 0 to 1. These different sulfate phases all influence the set behavior of portland cement. Because the procedures given in ASTM 1365 provide mass fraction data for all the major chemical constituents of portland cement, they also play an essential role in the validation of the cement phase volume fraction data.

Cement Phase Volume and Surface Area Fractions, Two-Point Correlation Functions

The most important data required by the VCCTL software for the cement being simulated are the volume and surface area fractions of the four major cement phases: alite, belite, aluminate, and ferrite. While volume fraction can be obtained via x-ray powder diffraction, acquisition of surface area fraction data is much more complex, involving scanning electron microscopy and multispectral image analysis. This process results in a false color, segmented image of a cross section of cement particles. The complete process of cement phase volume and surface area acquisition is detailed in Chapter 4.

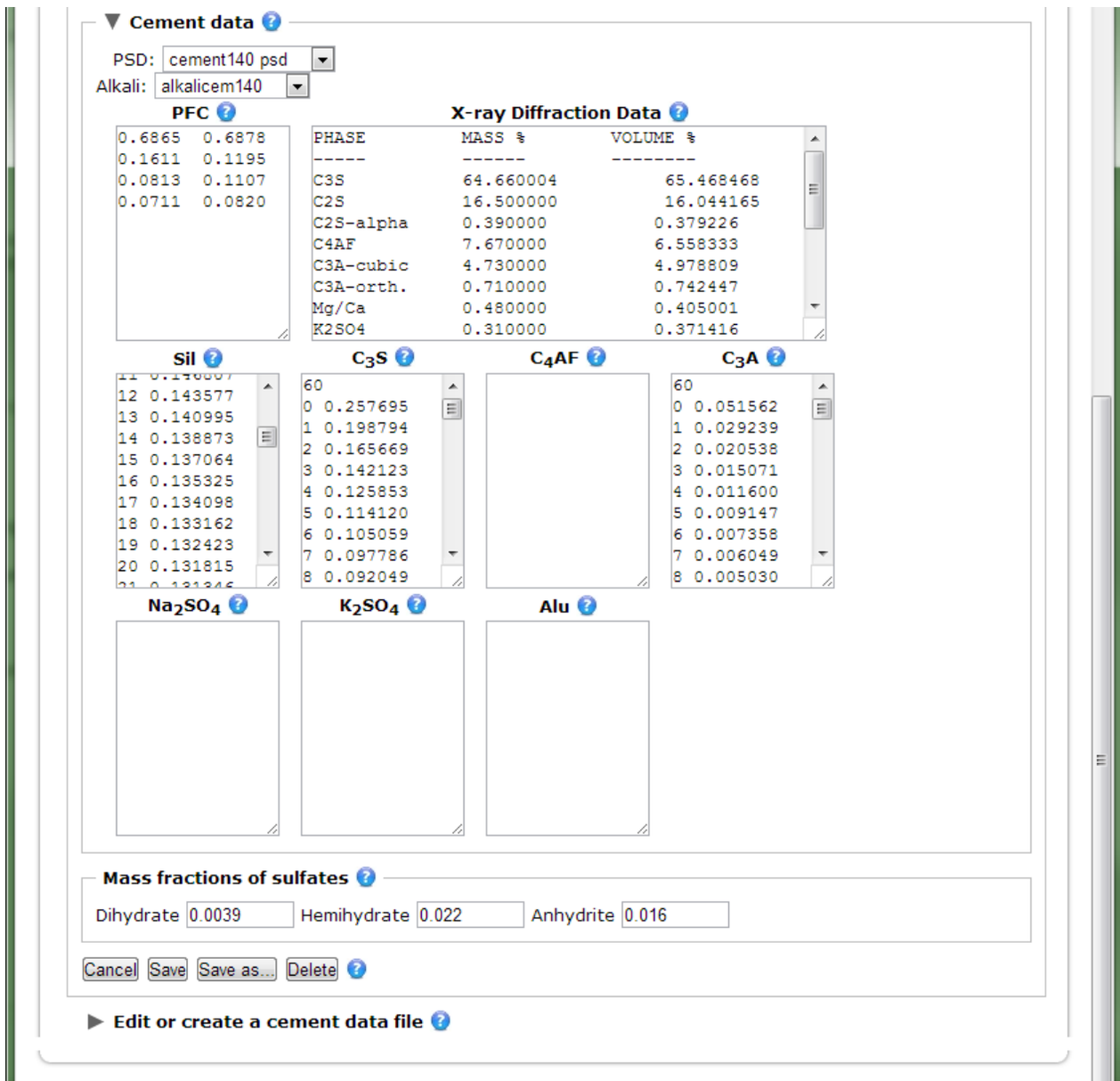


Figure 3-2: Cement Phase Data Input Screen

VCCTL software creates a digital microstructure based on data measured from a cement, and also calculates a spatial distribution of the cement particles as isotropic two-point correlation functions for each of the different phases. A two-point correlation function is a statistical representation of the correlation between random variables at different points in space. In this case, the correlation function for each phase describes the probability that a pixel a specific distance away from the current pixel is of the same phase as the current pixel. When the segmented image is measured to obtain phase fraction data, this data is also measured from the

image. The software uses these two-point correlation functions to build a three-dimensional microstructure that is consistent with these functions. Figure 3-2 details the screen in which the phase and volume fractions can be entered, as well as the two-point correlation functions for each phase.

Cement Particle Size Distribution

Laser particle size analysis is used to obtain the particle size distribution curve that the VCCTL software requires for cement. This information was obtained using a Horiba LA-950 Laser Particle Size Analyzer, which measures the diffraction of light scattered by particles as they pass through a laser beam. The machine uses compressed air to blow particles through the beam and sequentially measures the size of each particle. Particles are measured until the size distribution no longer changes. An example particle size distribution is shown in Figure 3-3.

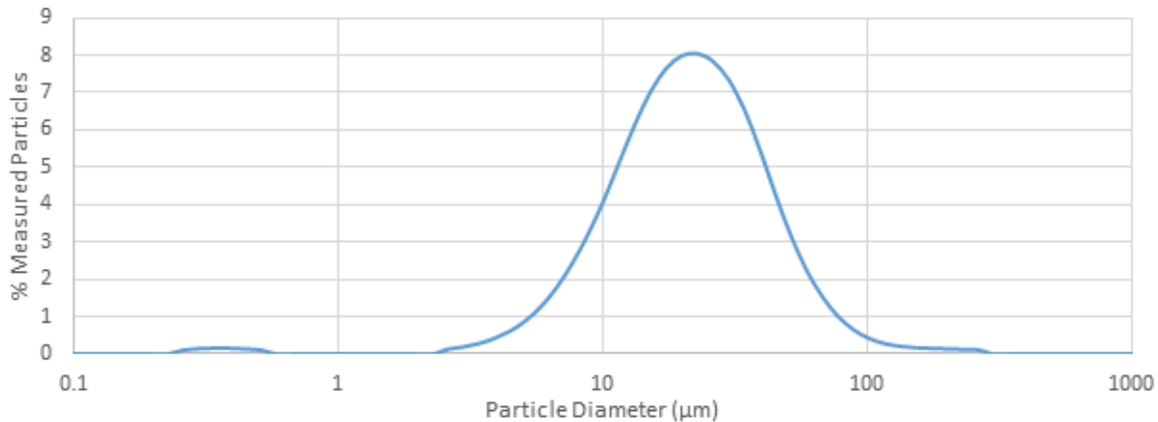


Figure 3-3: Cement Particle Size Distribution

Cement fineness is typically measured using the Blaine air-permeability test, which is detailed in ASTM C204. The test determines the fineness of portland cement based on the rate at which air under a pressure differential (via a manometer) passes through a cement powder bed. The rate at which the air passes through the cement is compared to the rate of a known standard, and the specific surface, in m^2/kg , is calculated via an empirical equation. While this method is useful for comparing the relative fineness of cements, it does not provide a particle size distribution. The VCCTL software requires a particle size distribution so that the digital microstructure will reflect the fineness of the selected cement. The cement fineness has a substantial impact on the rate of hydration, with finer cements resulting in higher rates of

hydration (Mindess & Young, 1981). This input is also selected on the screen shown in Figure 3-2.

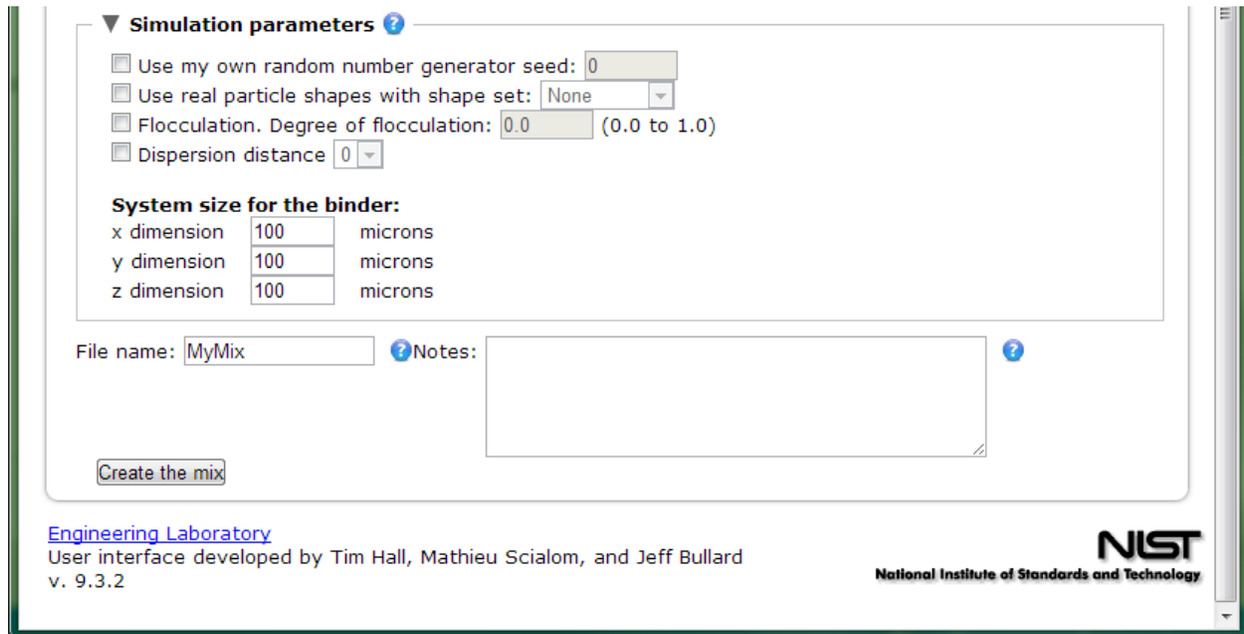


Figure 3-4: Microstructure Simulation Parameters

Cement Particle Shape Data

The VCCTL software builds the digital microstructure using spherical particles, though it also has provisions for the use of real cement particle shapes. These particle shapes are obtained using x-ray computed microtomography scans of real cement particles, which are then individually modeled using a spherical harmonic expansion of the particle surface. The coefficients of the equations used to model the particle can then be used to reconstruct the particle virtually with arbitrary size and orientation (Bullard & Garboczi, 2006). No cements were characterized in this way during this research; however, the software contains a library of cement particle shape sets. A cement from the library with particle shapes judged to be similar to that of the cement used in this research was used when simulations with real particle shapes were run. A screenshot showing this option is visible in Figure 3-4

Cement Particle Dispersion

The VCCTL software provides the option to create a simulation that considers dispersed cement particles as it composes the virtual microstructure. This option is provided to simulate the mutual repulsion of cement particles that occurs (Mamlouk & Zaniewski, 2010) when water-

reducing admixtures are present. The functional effect of this is that space between particles is more uniform, which can result in more accurate simulation of hydration and a corresponding increase in strength (Neville, 2011). When simulating mixes with water-reducing admixtures, this option (shown in Figure 3-4) was enabled and set to a value of one. The dispersion of particles can be set to 0, 1, or 2, with the values indicating the minimum number of pixels separating each particle in the virtual microstructure.

Binder System Size

By default, the volume of the microstructure created by the VCCTL software is 100 microns cubed. The size of the microstructure can be increased or decreased by the user, though a linear increase in the side of the virtual microstructure will result in a cubic increase in volume, impacting the time required for the computer modeling of microstructure operations.

Hydration Modeling Inputs

The following inputs are used by the VCCTL software to govern the behavior of the hydrating microstructure.

Isothermal Conduction Calorimetry – ASTM C1702

ASTM C1702 details a standard test method for the measurement of heat of hydration of hydraulic cementitious materials using isothermal conduction calorimetry, a process which has seen widespread adoption for cementitious research in the last few decades (Ferraro, 2009). The method details the construction and function of an isothermal conduction calorimeter, the recommended procedures for the preparation of samples, and explains the analysis of the results obtained from a calorimeter to determine heat of hydration. The test method defines an isothermal conduction calorimeter as a constant temperature heat sink to which two heat flux sensors and sample holders are attached in a manner that insures good thermal conductivity. The two sample holders are required for the analysis of one sample, one for the cementitious sample and one for a reference sample that evolves no heat.

The heat of hydration evolved by the sample flows across the heat sensor and into the heat sink. The measured heat of hydration is the difference between the heat flow of the sample cell and the reference cell. A schematic diagram of a basic isothermal calorimeter is shown in Figure 3-5 (ASTM C1702, 2009).

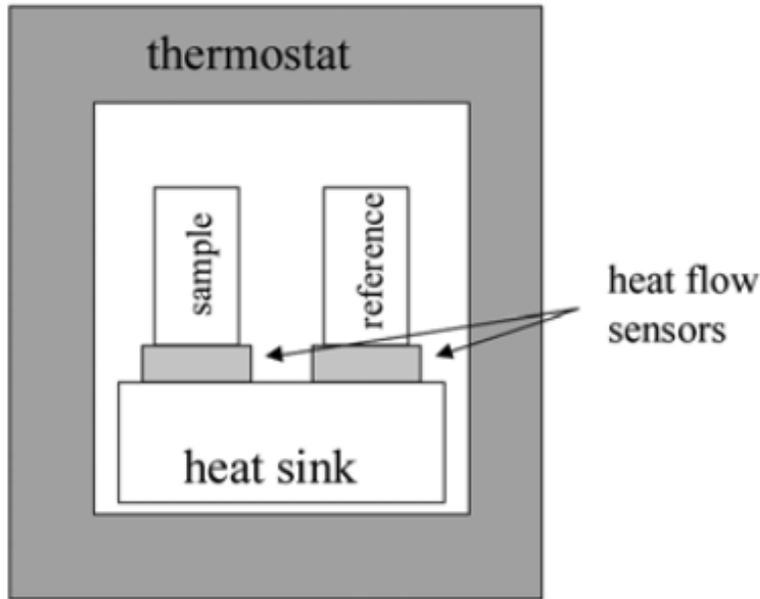


Figure 3-5: Simple Isothermal Calorimeter (ASTM C1702, 2009).

One of the defining characteristics of hydraulic cements is that they exhibit an exothermic reaction when mixed with water. Ideally, the heat released from this reaction should be measured from the moment the sample is mixed with water and sample the should be in thermal equilibrium with the environment within the calorimeter, as any heat flow through the sensor that exists prior to hydration can distort the measurements acquired. The standardized test method details a procedure by which these two conditions can be met and is referred to as internal mixing.

An admix ampoule, shown mounted onto a sample vial in Figure 3-6, is used to facilitate internal mixing. The admix ampoule has an o-ring seal, syringes that hold the mix water, and a mechanical stirrer for mixing. The admix ampoule-sample vial assembly is placed inside the isothermal calorimeter and allowed to reach thermal equilibrium prior to mixing. After equilibrium is reached, the water is ejected from the syringes into the sample vial containing the cementitious material, and the paste is mixed using the manual stirrer. The total heat evolved is typically recorded for 7 days.

The specification also outlines an alternative method for sample mixing, in which the sample is mixed outside the calorimeter and immediately inserted and measured. While much simpler than the previous procedure to execute, the test specification acknowledges a few

shortcomings of this method (ASTM C1702, 2009). The mixing of the specimen prior to insertion in the calorimeter necessarily means that the earliest stages of hydration are missed and that the sample is not at thermal equilibrium when the test is started.



Figure 3-6: Admix Ampoule

The data acquired before the sample reaches equilibrium has to be corrected, the procedure for doing so is detailed in the specification. For this research, the internal mixing procedure was chosen because the earliest stages of hydration were considered important for

evaluating the effects of admixtures, the acknowledged shortcomings of the external mixing procedure in the test specification, and past research showing the viability of the internal mixing procedure (Ferraro, 2009).

The specific sample preparation procedure used for isothermal calorimetry analysis was as follows. The quantities of cement and water to be used were calculated such that the total heat capacity of cement and water together were equal to that of the reference (inert) specimen. The cement was then carefully introduced to the sample vial, while monitoring the mass on a research balance, until the desired mass was reached. Deionized water was carefully drawn into the admix ampoule and measured on the balance. The process was repeated until the desired mass of water was contained in the syringes. Afterwards, a small amount of air was drawn into the tips of the syringes to help prevent drops of water from leaking onto the cementitious material while the sample was reaching thermal equilibrium within the isothermal calorimeter. Vials and admix ampoules were assembled with a small amount of vacuum grease to seal connections and then inserted into the calorimeter. At least three companion specimens for each mixture were used, with two mixes measured during one 7-day test period.



Figure 3-7: Tam Air Isothermal Calorimeter with Sample and Admix Ampoule Loaded

For this research a TAM Air isothermal calorimeter was used, shown in Figure 3-7. The calorimeter consisted of 8 channels, each channel consisting of two cells in the configuration shown in Figure 3-7. Samples were allowed to equilibrate for at least 12 hours overnight. Measurements began for each sample when the water was injected into the sample vial. Samples were stirred manually for 60 seconds each, after which measurements were taken for seven days.

Curing conditions

Thermal

Conditions:

- isothermal
- semi-adiabatic
- adiabatic

Initial temperature: °C

▶ Aggregate

Aging

Hydrate for days

... Or stop at degree of hydration:

- Use time conversion
 - Time conversion factor h/cycle²
- Use a calorimetry file
- Use a chemical shrinkage file

Saturation conditions

- saturated
- sealed

▶ **Simulation parameters**

▼ **Data output**

Evaluate percolation of porosity every: hours

Evaluate percolation of total solids every: hours

Evaluate individual particle hydration every: hours

- Output hydrating microstructure every hours
- Specify file with times for output
- Output hydration movie frame every hours

File name:

Figure 3-8: Hydration Simulation Input Parameters

The resulting power curve for the seven-day hydration reaction can be integrated over time to obtain a heat of hydration curve, which in turn can be input directly into the VCCTL

software as a text file. The input screen for the selection of a heat of hydration curve can be seen in Figure 3-8. For each computational cycle of the hydration reaction, the software calculates the amount of heat released. Heat of hydration data is not required by the software for hydration simulations, as the model assigns the same length (user input) to each cycle; however, if the experimentally measured heat of hydration data (via ASTM C1702) is entered into the software, the model will assign a length to each computational cycle of the simulation such that the heat evolved during that cycle matches up with the input heat of hydration curve.

Curing Conditions

The VCCTL software provides input options to simulate different possible curing conditions for the hydrating microstructure, shown in Figure 3-8. The “Thermal” condition options regulate the rate at which the heat evolved during hydration leaves the microstructure. The “Isothermal” thermal condition selection sets the rate to be equal to the rate at which heat is evolved (keeping the microstructure at ambient temperature), the “Semi-Adiabatic” option sets the rate at a user-specified value, and the “Adiabatic” option sets the rate such that no heat leaves the microstructure.

The options for “Saturation conditions” influence the water available for the hydration simulation. The saturated option allows the microstructure to never run out of water for hydration reactions, while the sealed option limits the available water to that which is in the initial microstructure. For the purposes of this research, the saturated option was always selected.

Data Output

The VCCTL allows the user to specify the intervals at which specific types of data are measured from the hydrating microstructure, including the degree to which the pore space in the microstructure is interconnected (percolation of porosity), the degree to which the solids in the microstructure are interconnected (percolation of total solids) and the degree to which each particle within the microstructure is hydrated. These options are shown in Figure 3-8, and are computed at user specified intervals. Also part of the data output section are the intervals at which to output microstructures for moduli calculations.

Aggregate Input Data

Most of the input data required by the VCCTL software for aggregates come from ASTM standard tests that are performed on aggregates as a matter of course, the results of which are readily available. A brief summary of these tests follows:

- ASTM C136 – Standard Test Method for the Sieve Analysis of Fine and Coarse Aggregate. This test is run on all aggregates used at the FDOT;
- ASTM C127 – Standard Test Method for Density, Relative Density (Specific Gravity), and Absorption of Coarse Aggregate. This test is also run for all aggregates used at the FDOT.

The primary use of the aggregate input data by the software is simply to convert the mass fractions to volume fractions when batching the virtual mix. Aggregate bulk and shear moduli are not among the data readily available to characterize aggregate. While the aggregate properties can be measured from rock cores sourced from the mine where the aggregate originated, there is a degree of inherent variability to natural aggregate that makes obtaining representative cores difficult. This process was not performed for this research due to time and scope constraints. Instead, some values were assumed for the coarse and fine aggregate used in this study based on the knowledge that the VCCTL software requires this data for the calculation of the modulus of concrete. As concrete is mostly aggregate by volume, bulk and shear moduli potentially have a substantial influence on the results. Thus, future research should include a detailed investigation of these properties.

The VCCTL software accounts for aggregate shape data as a statistical distribution of spherical harmonic coefficients, which is obtained via x-ray computed tomography. The software includes shape data for a variety of aggregates in its built-in library of materials, and the influence of aggregate shape on simulation results was investigated as part of this study and is discussed in Chapter 7.

CHAPTER 4 SEM MICROANALYSIS

Overview

The phase volume fractions and surface area fractions are required inputs for VCCTL software to simulate cement microstructure as it hydrates, and are acquired via a combination of scanning electron microscopy and image analysis. The process was originally developed at NIST and consists of three primary operations.

1. Backscatter Electron and X-Ray Map Image Acquisition,
2. Image Processing, and
3. Creation and Analysis of Segmented Image.

Before any of the data acquisition operations can take place a significant amount of sample preparation is required to enable the imaging of the cement being analyzed within the scanning electron microscope.

Sample Preparation

Procedures Developed by NIST

The phase volume and surface area fraction data required by the VCCTL software is obtained through a complex process involving image acquisition using a scanning electron microscope (SEM). Before this process can even begin however, the cement being analyzed must be prepared so that imaging is possible. The analysis process examines cross sections of cement grains which must be flat and very smooth, free of scratches, tear outs, and other imperfections that could skew measurement or cause imaging artifacts. The process required to achieve this is complex and sensitive. The original procedure for the preparation of cement powder specimens for imaging in this way was developed by NIST to provide inputs for different cement hydration models. This original process, outlined in the next few paragraphs, was used as a starting point for the procedure developed for this project.

The cement is first mixed with an optical grade epoxy resin in proportions that result in a stiff paste. The paste is then formed into a ball and pressed into a specimen mold. The paste is consolidated by tapping the mold sharply upon the table top. Once consolidation is complete, the specimen is then cured according to the guidelines recommended by the epoxy manufacturer.

After the specimen is cured, a fresh surface is exposed by grinding or sectioning with a wafering saw.

Following the resin encapsulation process, the specimen is ground with silicon carbide sandpaper using progressively finer abrasives, stopping at 600 grit. This process serves to remove the damage to the surface left by the diamond saw blade, and prepares the surface for the final polishing steps. The specimen is polished using progressively finer grit diamond abrasive and a polishing cloth, specifically 6, 3, 1, and 0.25 μm . At each stage the specimen is polished until the imperfections left from the preceding abrasive are removed. Ethanol-based lubricants are used throughout the cutting, grinding, and polishing process in order to prevent further hydration of the cement grains or paste. The final polished surface is carefully cleaned, after which a carbon coating is applied to render the sample conductive for imaging in the SEM.

Improvements to Sample Preparation

The original procedure developed by NIST was almost 20 years old when this research project began. Though the fundamental process remains the same, substantial improvements to the speed of preparation and consistency of quality were made. One of the biggest improvements made to the existing procedure was the automation of the grinding and polishing operations. The original process involved hand grinding of samples using silicon carbide paper on plates of glass, followed by hand polishing using a manual lapping wheel. These operations are very labor intensive and require a considerable degree of operator skill.



Figure 4-1: Saphir 550 Semi Automated Grinder Polisher

An automated grinder polisher was acquired for this project specifically for the purpose of improving sample quality and reducing preparation time. Shown in Figure 4-1, the Saphir 550 grinder polisher allows for precise control of all the variables involved in the process of grinding and polishing samples. The basic operation of a polisher is simple. The polisher consists of a working wheel with some type of abrasive, either a silicon carbide paper, a resin bonded diamond disc, or a polishing cloth with a diamond suspension, and a powerhead that controls the motion and pressure applied to the sample holder, based on user preferences. The specimen to be polished is then held against the working wheel as it spins and moves in a circular motion. In the case of a manual grinder polisher, the user would be in charge of holding the specimen against the working wheel. This semi-automated grinder polisher, however, has a powerhead that can accommodate sample holders in which up to six two-inch specimens can be mounted. The powerhead can spin at a user specified RPM, ranging from 30 to 150 rpm, and the working wheel can spin from 50 to 450 rpm.

When grinding and polishing manually, the user is responsible for the amount of force with which the specimen is pressed against the working wheel, whereas the Saphir 550 allows each of up to six specimens to be pressed against the working wheel with a user specified force

ranging from 5-100 N. In addition, the user can specify a two-stage force profile, wherein the force applied is one value for an initial period of time, ramps up to a higher value for the majority of the operation, and then falls back off to the first value for the final stage of the operation. This allows for a period of lighter material removal at the end of a grinding or polishing stage, resulting in a better surface finish. The Saphir 550 maintains the surface flatness and uniformity of the specimens through the use of an automated powerhead and individual specimen forces.

A typical grinding and polishing operation consists of several stages, using progressively finer abrasives in each stage. Depending on the initial quality of the specimen's surface finish, a number of grinding operations may be used. When the surface has been ground to an acceptable finish, the polishing operations commence. With the use of a polishing cloth and diamond abrasive, the specimen is polished using progressively finer abrasives until the surface finish reaches the desired level uniformity. The user is responsible for all aspects of the operation, including the application of lubricant to the working wheel, determination of when to switch to the next abrasive stage, how much abrasive to use, and how much force to apply during each stage. The advantage of a semi-automated grinder polisher is that once the optimum series of grinding and polishing operations have been determined, they can be programmed into the grinder polisher. The Saphir 550 will perform all of the tasks required during a grinding and polishing operation except for the changing of the working wheel when switching to a finer abrasive.

During the development of the grinding and polishing procedure, inspection of the surface finish of the specimen after each step was useful to determine its efficacy. An optical microscope, a Nikon AZ-100 shown in Figure 4-2, was purchased as part of this project and was used for this purpose.. With a maximum resolution of about 1.5 microns and a magnification of up to 650x, this microscope allowed for the evaluation of the flatness and surface quality of samples after each polishing step.



Figure 4-2: Optical Microscope

Some experimentation was necessary to develop a grinding and polishing process that resulted in specimens of suitable quality as there were issues with cement particles being torn from the resin during the grinding steps at the time of initial trials. This problem was overcome by changing the initial grinding media, as well as by switching to a harder resin. A resin-bonded diamond abrasive disk was used for the first grinding steps initially, but it was found that the hardness of the diamond abrasive created relatively deep scratches in the specimens that could not be removed during the later polishing steps. The solution to this was to switch to silicon carbide grinding papers which offered a more progressive grind, as the softer silicon carbide actually became finer in grit as the abrasive eroded in the polishing process. Other variables in the grinding and polishing process which were investigated included, varying the specimen force, lubricant and abrasive dispensing intervals, and rotation speeds. Table 4-1 is a summary of the procedures used for specimen preparation in this research.

Table 4-1: Grinding and Polishing Procedure

Step	Time	Abrasive	Abrasive Type	Media	Force	Head Speed (rpm)	Wheel Speed (rpm)	Lubricant Interval	Abrasive Interval
1	1:00	600 grit	Silicon Carbide	Grinding Paper	20 N	150	149	0:30	-
2	1:00	1200 grit	Silicon Carbide	Grinding Paper	20 N	150	149	0:30	-
3	1:00	4000 grit	Silicon Carbide	Grinding Paper	20 N	150	149	0:30	-
4	2:00	6 μm	Diamond Suspension	Silk Cloth	15 N	150	149	0:45	0:45
5	2:00	3 μm	Diamond Suspension	Silk Cloth	15 N	150	149	0:45	0:45
6	2:00	1 μm	Diamond Suspension	Silk Cloth	15 N	150	149	0:45	0:45
7	4:00	0.25 μm	Fumed Silica	Silk Cloth	10 N	150	149	0:30	2:00

The final step to specimen preparation was to deposit a conductive coating on the surface of the sample which was performed using an evaporative carbon coater, shown in Figure 4-3. The Cressington 108carbon/A carbon coater used a high electrical current combined with a sharpened graphite rod to coat a specimen in a nanometer-thick layer of elemental carbon. The specimen was placed in the coating chamber, the chamber was evacuated using a rotary pump, and current was then passed through the carbon rod, causing the carbon to evaporate and then deposit on the specimen. Specimens were coated for 4 seconds, three times, which was found to produce a coating thickness that enabled effective imaging (good signal with no charge buildup).



Figure 4-3: Evaporative Carbon Coater

Backscattered Electron and X-Ray Map Image Acquisition

After the completion of the sample preparation process, an unhydrated cement sample was analyzed using a scanning electron microscope. The two primary imaging modes used in this process were backscattered electron (BSE) and energy dispersive x-ray (EDS) mapping. The BSE mode created an image wherein the contrast was determined by the relative average atomic number (Z) of the different phases, with the brightest phases having the highest average atomic number. A typical BSE image is presented in Figure 4-4. The brightest phase present is the ferrite phase (C_4AF), due to the presence of iron, with is denser relative to the other phases, which are composed of mainly calcium, aluminum, and silicon. The darkest phase is the epoxy substrate, as it is a plastic and thus composed primarily of relatively low-atomic number carbon polymer chains. The phases in order of darkest to lightest (lowest to highest atomic number) are the voids (epoxy-filled), gypsum, C_3A , C_2S , C_3S , and C_4AF .

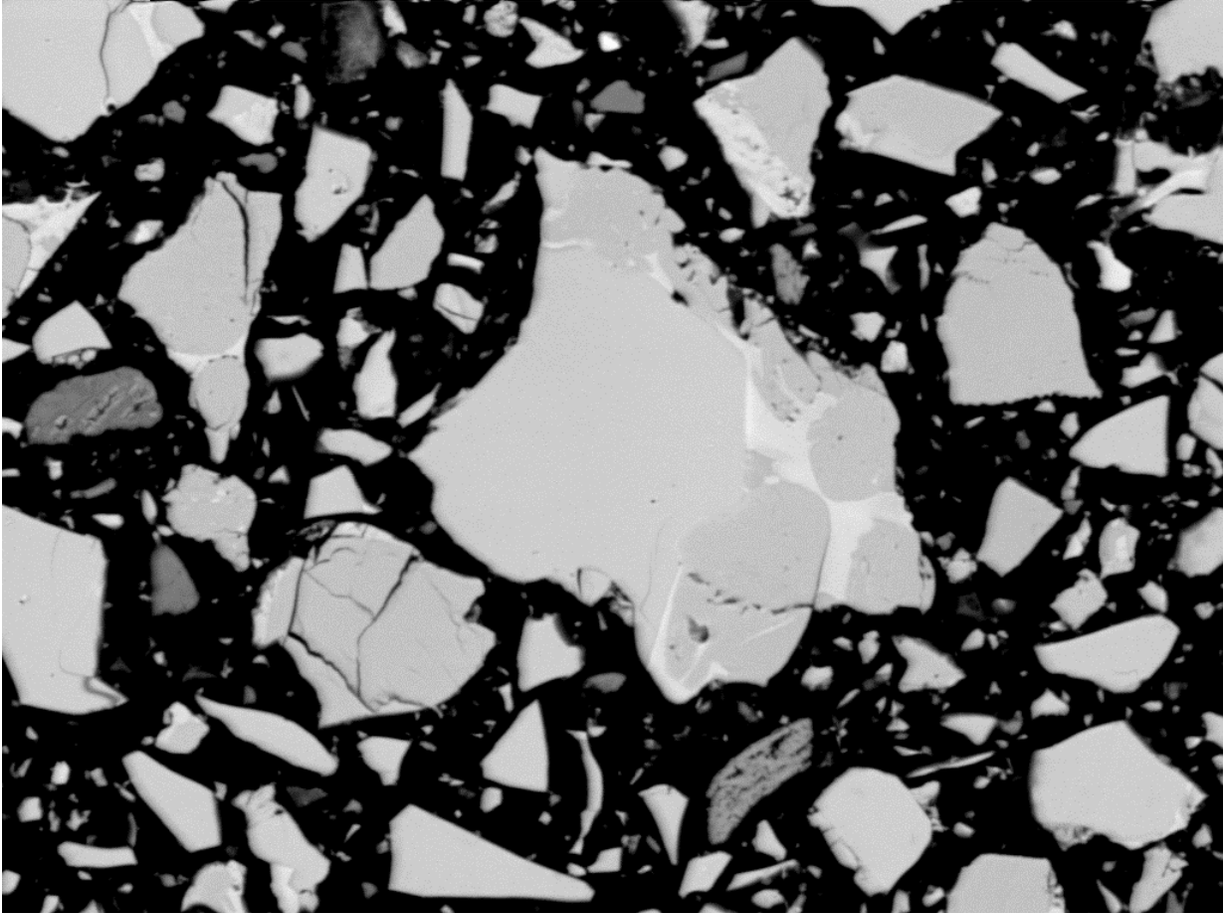


Figure 4-4: Cement Grain Close-up at 1600x magnification

Figure 4-4 is a close-up of a polished cement grain. It is possible to discern the four primary cement phases with a reasonable degree of accuracy based on the relative grey levels and the morphology of the phases (aluminate is typically dispersed within ferrite, for instance). However, computerized quantitative analysis of the area and perimeter of each phase was not possible without more information, as different phases (belite and aluminite) had overlapping ranges of grey values. BSE images used for analysis were typically of a larger area than that shown in Figure 4-4. An example of a typical image used for this process is provided in Figure 4-5.

The second imaging mode needed for this technique, EDS, was used to create an image that showed the location and relative concentration of specific elements within the cement sample. Elements can be identified due to the characteristic nature of the electron beam-

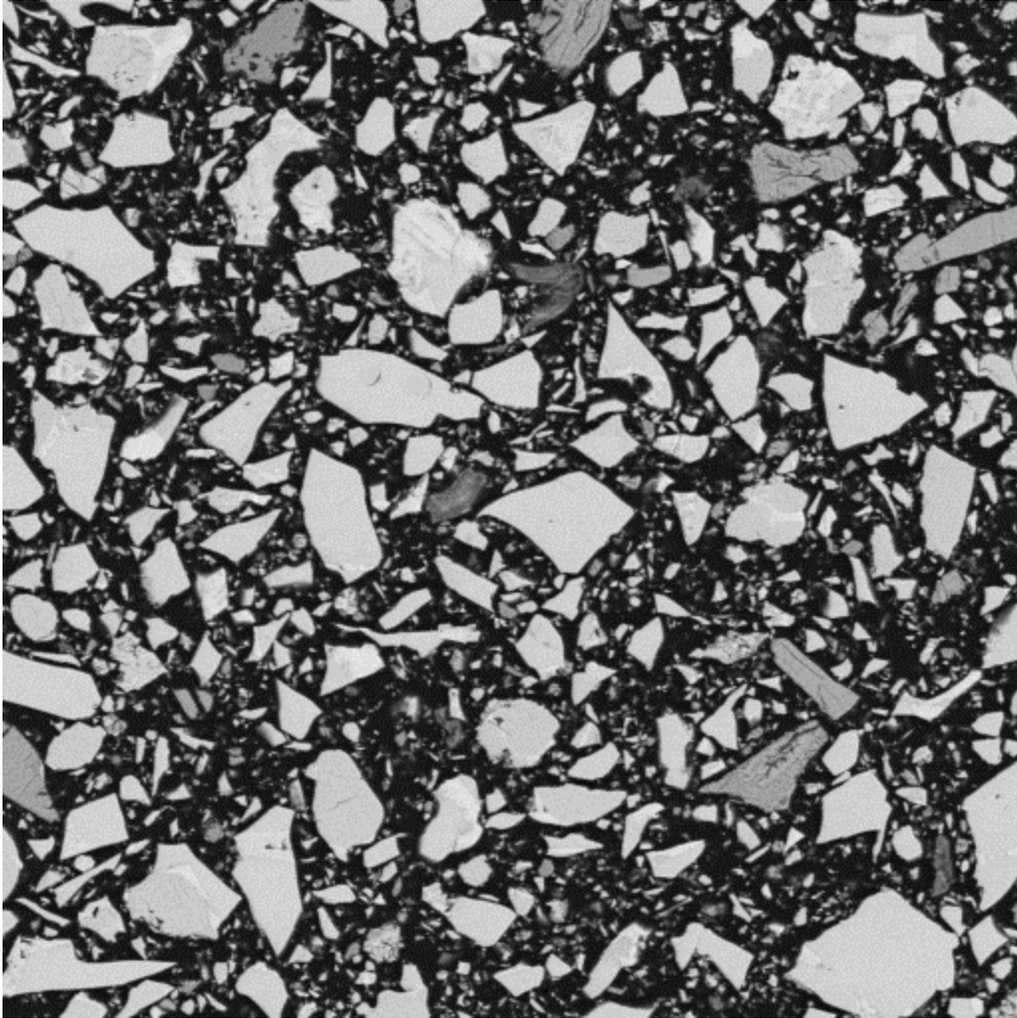


Figure 4-5: BSE Image as Used for Phase Analysis, 800x Magnification

specimen interactions, where the number of interactions for a particular element determines the relative brightness (elemental contrast) in a backscattered SEM image, and is related to the atomic number of the element. The interaction of the electron beam with the electron clouds of atoms in the specimen results in the ejection of inner-shell electrons which result in higher-energy outer-shell electrons moving to empty lower-energy inner shells, which results in the release of x-ray photons. The photons have energies equal to the differences in the energy states of the donor and recipient orbitals in the affected atoms. These photon energies uniquely identify the corresponding elements and are referred to as characteristic energies. By scanning a selected area of the specimen for x-ray photons of a specific energy, a “map” is created for the particular element identified by the characteristic energy.

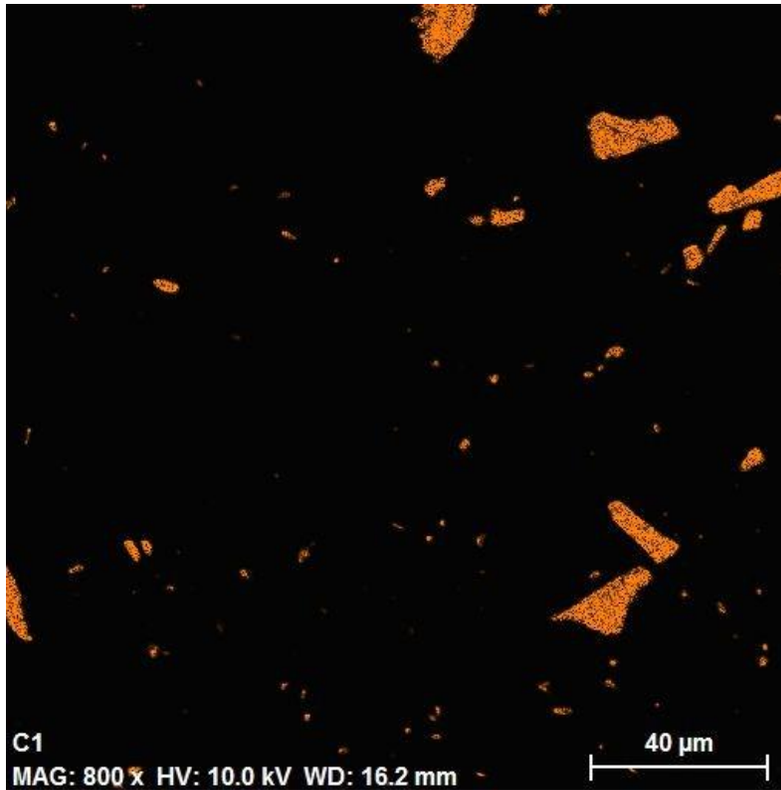


Figure 4-6: Elemental Map for Sulfur

An example of an EDS elemental map of the region shown in Figure 4-5 is presented in Figure 4-6. The areas that are not black denote the presence of sulfur. This sulfur map and corresponding maps for aluminum, sodium, and potassium, provide the additional information needed to distinguish the phases present in the BSE image that would otherwise be partially indistinguishable due to overlapping grey values.

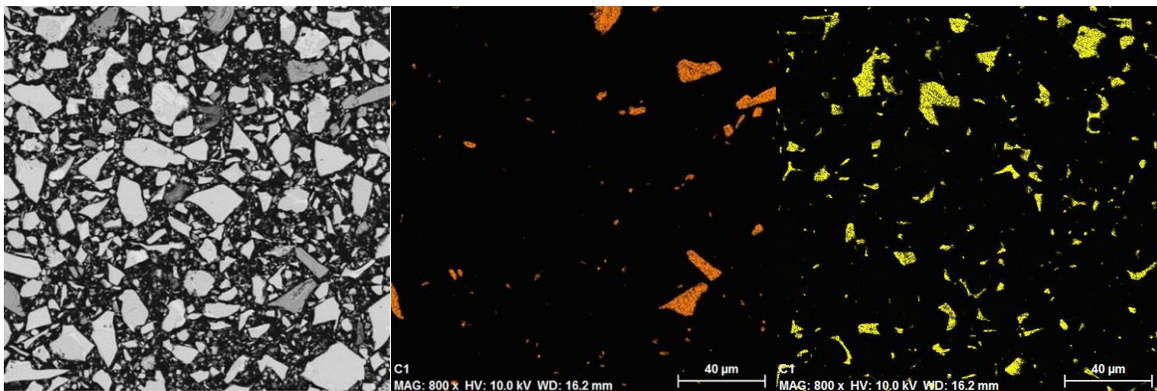


Figure 4-7: BSE, Sulfur, and Aluminum Maps Required to Distinguish Alite, Belite, Aluminate, Ferrite, and Gypsum

The images shown in Figure 4-7 are those required to distinguish the 4 primary phases in cement as well as gypsum. They are, from left to right, the BSE image, the sulfur map, and the aluminum map.

Image Processing

Multispec is a program that has been developed at Purdue University for the purpose of processing and analyzing multispectral and hyperspectral images and was used in this research to combine images such as those shown in Figure 4-7. A multispectral image is created by combining different imaging modes of the same area. Typical multispectral images of the same object could be some combination of thermal imaging, visual imaging, and infrared imaging. By assigning the BSE image and each EDS map to a specific color channel (red, green, blue) a multispectral image of the cement powder is created. This multispectral image contains more information on phase composition than any of the individual images that it was created from. Figure 4-8 illustrates that by assigning the BSE image to the red channel, the aluminum map to the blue channel, and the sulfur map to the green channel, it is easier to discern the different phases of portland cement in comparison to the grayscale image created by BSE imaging (Figure 4-4). Multispec has the capability to automatically analyze and assign a specific phase value to each pixel in the image using user defined training fields. The fields are visible in Figure 4-8, and are regions selected by the analyst that contain a specific phase. These user-selected regions provide a reference for the program which it can apply in the classification of the entire image.

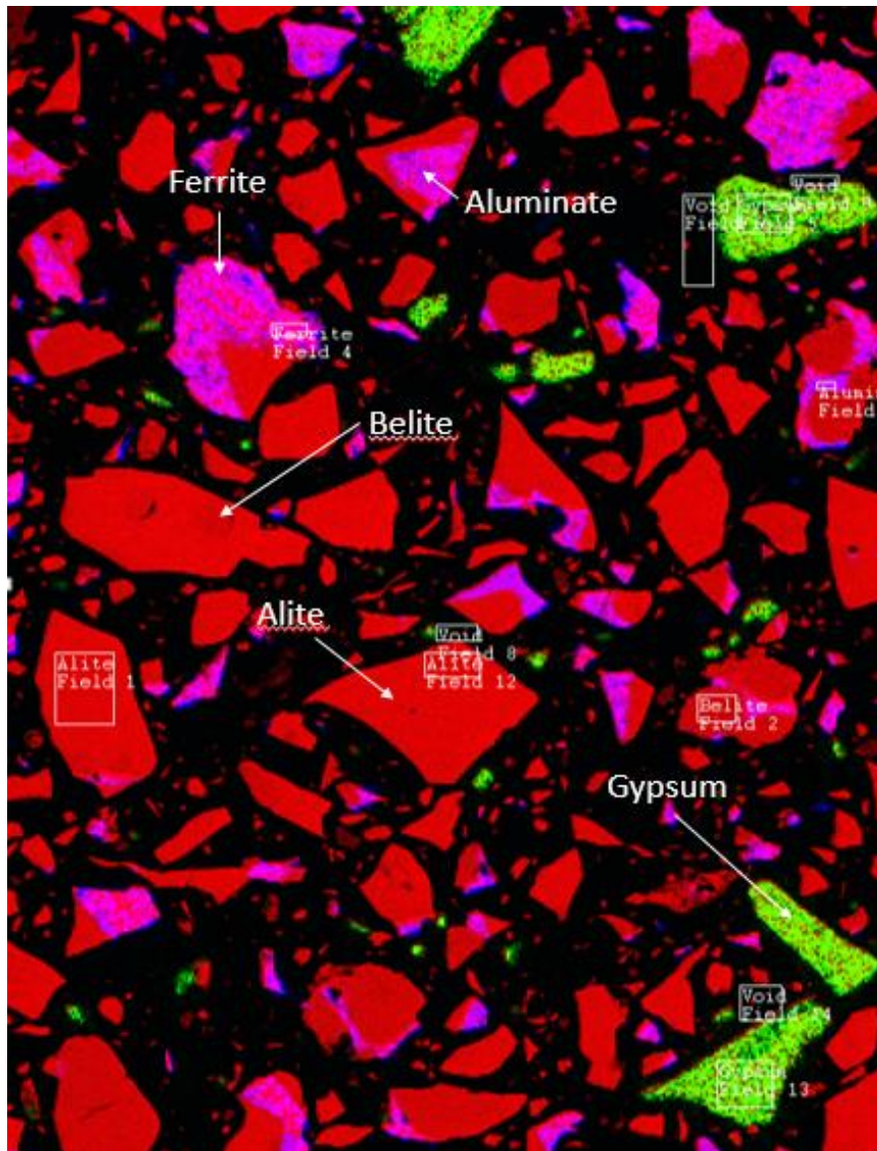


Figure 4-8: Phase Classification Using Multispectral Image and User-Defined Training Fields

The selection of these training fields is critical to the proper classification of the phases, and requires substantial practice. Once training fields have been selected, the program classifies the image, using the settings displayed in Figure 4-9. The result is then saved to a .GIS file.

A portion of the results provided by the Multispec program includes a statistical analysis and the calculation of the accuracy of the classification. The statistical result is based on the reference fields and the reliability of the classification, as well as the distribution of each of the phases being analyzed. These results for this image set are shown in Figure 4-10.

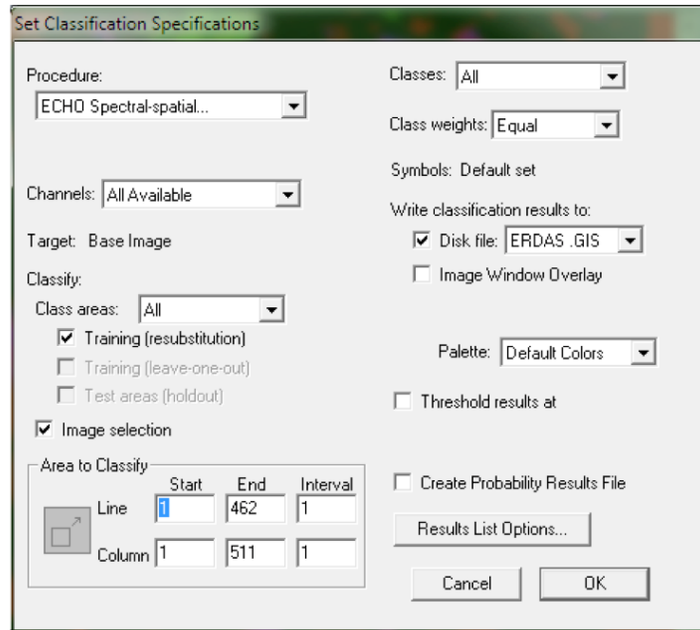


Figure 4-9: Image Classification Dialog

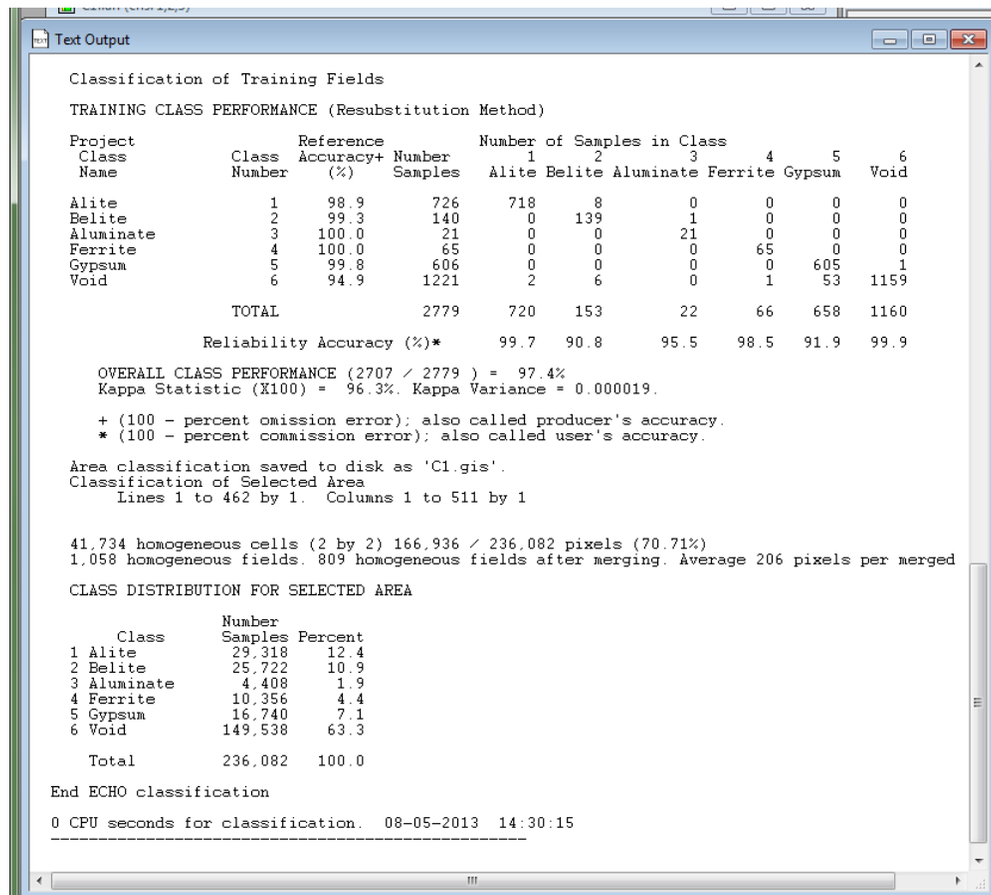


Figure 4-10: Statistical Analysis of Classification

Creation of Segmented Image

When the selection of fields and classification of phases has been performed such that it accurately represents the analyst's interpretation of the BSE electron image, the resulting .GIS file is viewed. The result is a false color image in which each color represents a different phase. Figure 4-11 shows the image resulting from the classification of the field set being used in this example. The image is very close to the final segmented image, and represents the end of the segmentation process itself.

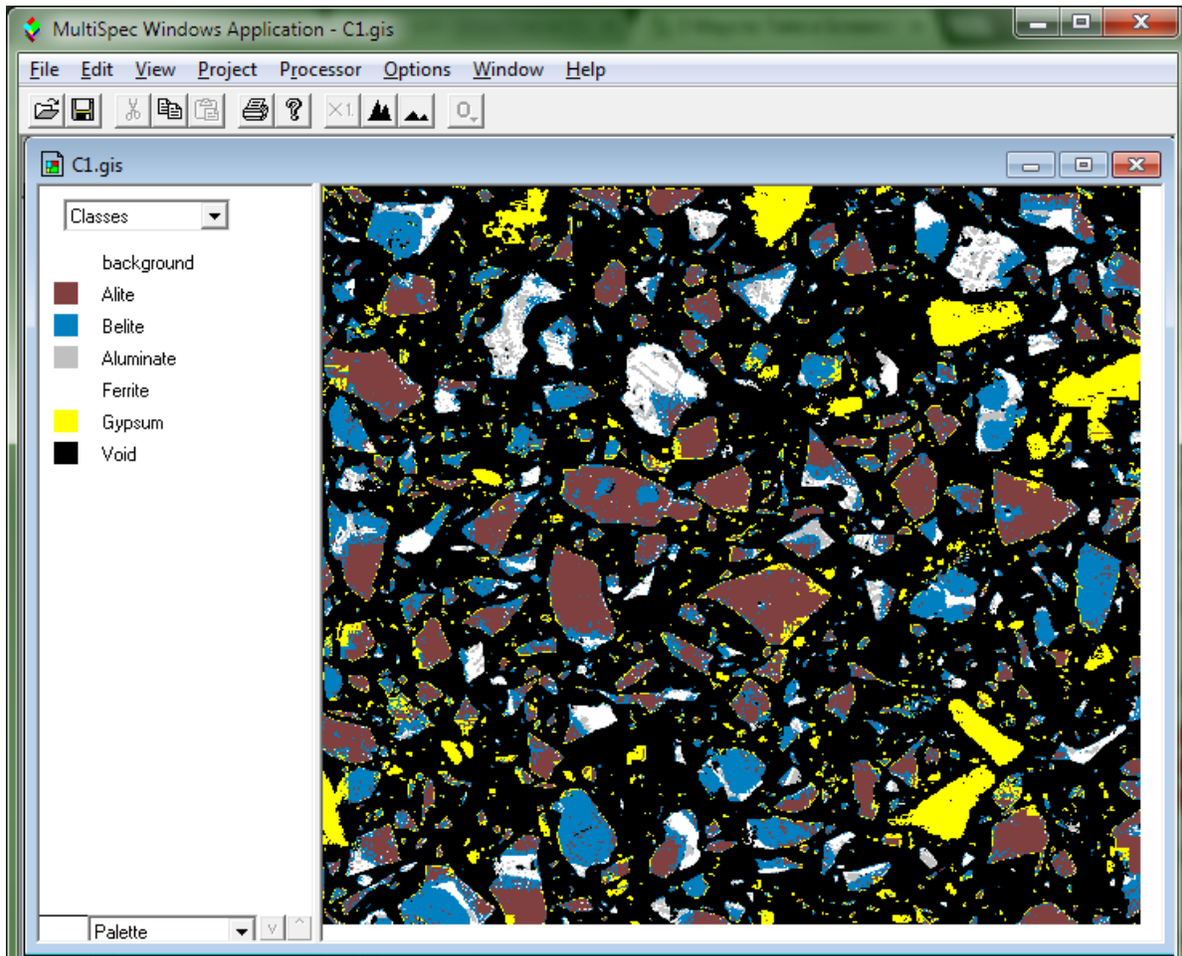


Figure 4-11: Segmented Image

In order to take measurements of the segmented image, some post processing is required. For this process, a program called ImageJ is used to perform a "Thresholded Blur" operation. This operation looks at a pixel and compares it to all the pixels within a user defined surrounding radius. This algorithm allows for the removal of fine noise without distorting or blurring the

boundaries between different regions. This is critical as the surface area fraction measurement can be skewed by excessive noise in the segmented image.

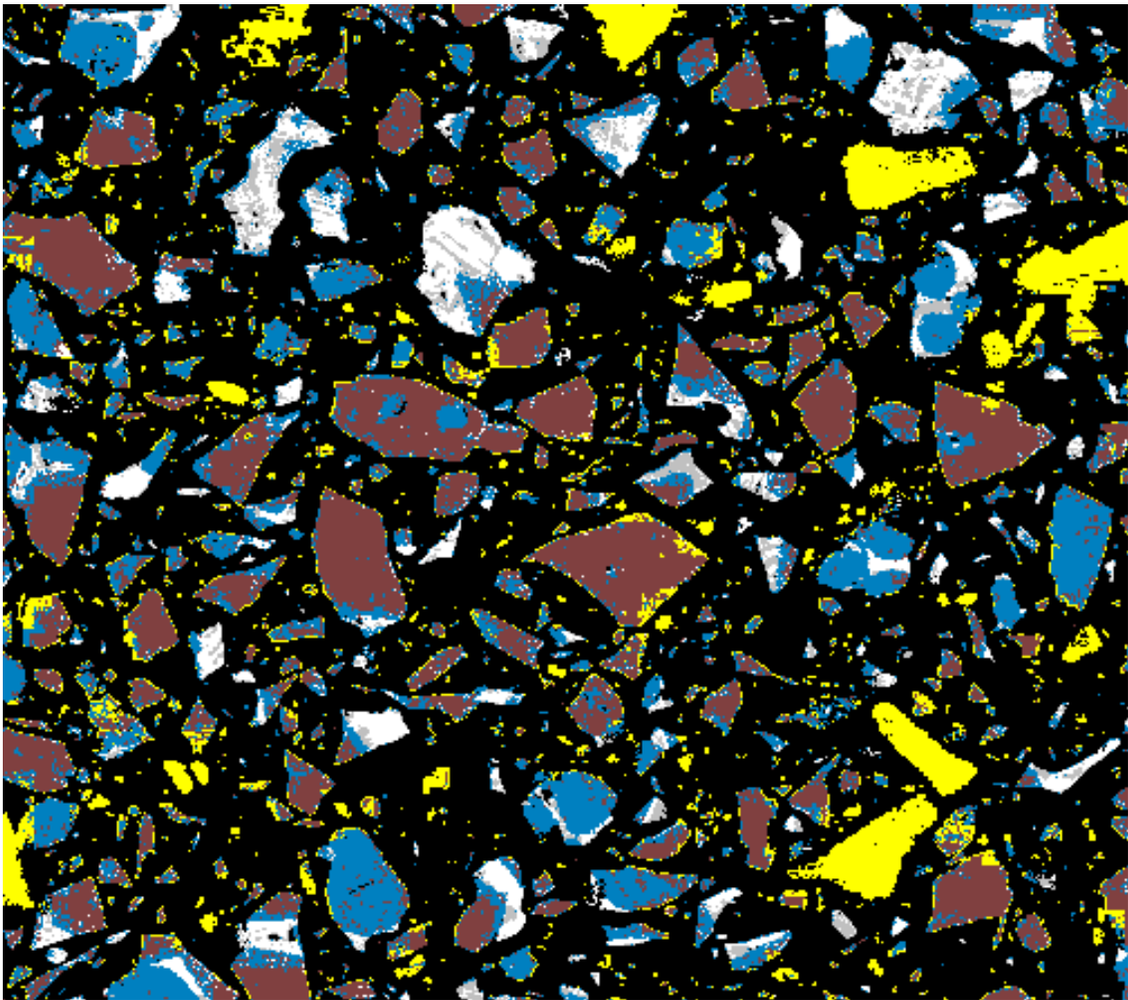


Figure 4-12: Segmented Image Prior to Thresholded Blur Operation

Figure 4-12 shows the image prior to the application of the thresholded blur filter. The settings for the filter are varied and the image examined, using the preview function, to remove the noise without removing an excessive amount of detail from the image. The only value that must remain the same is the threshold value, which is held at 1.

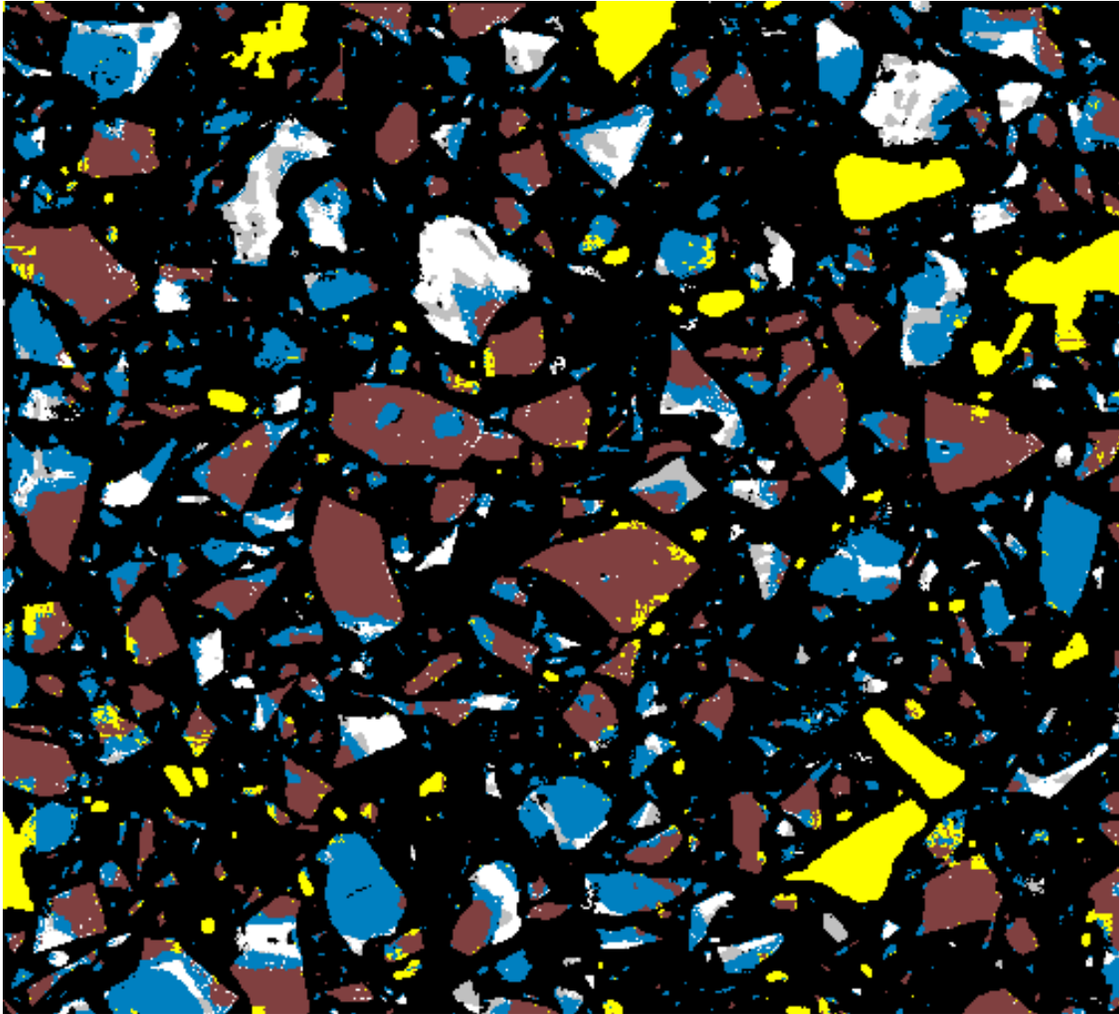


Figure 4-13: Segmented Image after Blur

Figure 4-13 shows the result of the thresholded blur which clearly displays a substantial reduction in noise and a general improvement of the distinctions between the different phases, and is a typical resultant image from which area and perimeter measurements of the portland cement specimen are taken. The image is saved in “ImageJ” as an indexed text image, in which each pixel is represented by a numeric value corresponding to the chemical phase of that pixel. This image is then used as the input for an executable provided by NIST that measures the volume and surface area fraction of each phase, and also determines the relative spatial correlations of each phase. Volume and surface area fractions are measured as area and perimeter fractions, since for isotropic systems (in which no preferential orientation is observed) the two parameters are the same (Bentz, 2002). The same is true for the spatial correlation functions. The

program provides an output as a set of files that can be archived as a .zip file and used for input directly into the VCCTL software for analysis.

One aspect of this process that is important to consider is that each image set created represents what is a very small cross section of a larger bulk material. In order to obtain statistically valid data for input into the VCCTL software, the process must be repeated for many image sets. The results of each image set are then averaged to obtain a more representative dataset. The following summarize the results of the analysis of 14 fields, including the above example, and the field analysis data are provided in Appendix B.

Table 4-2: Volume and Surface Area Fraction Averages

Phase	Volume %	SA %
C ₃ S	50.72	32.43
C ₂ S	27.94	44.04
C ₃ A	8.620	13.73
C ₄ AF	12.71	9.781

Table 4-3: Standard Deviations of Volume and Surface Area Fraction Measurements

Phase	Volume %	SA %
C ₃ S	6.62	6.92
C ₂ S	4.57	6.48
C ₃ A	3.81	7.20
C ₄ AF	3.59	4.88

The results presented in Table 4-2 provide a summary of volume and surface area fractions for the cement used in this study. The data displayed in Table 4-3 provide a summary of the standard deviation of the surface area and volume fractions for each phase. The range of standard deviation of the data obtained for the surface area fraction for each phase was between 3.0 and 7.5%. Some degree of variability between the fields is to be expected since the fields are not large enough to insure that they provide a representative sample for analysis. However, since the specimen processing and image preparation are rigorous, it is possible that the process can be refined further with better sample preparation. It is expected that the variation in the phase

acquisition from image analysis of SEM EDS data can be reduced by acquiring a better final polish of the samples analyzed.

This phase composition data can also be compared to that obtained from x-ray powder diffraction data using Rietveld analysis. The refinement of this process is an ongoing effort, and as they are further refined, the simulations should more closely represent the actual concretes being simulated.

Automated Cement Characterization

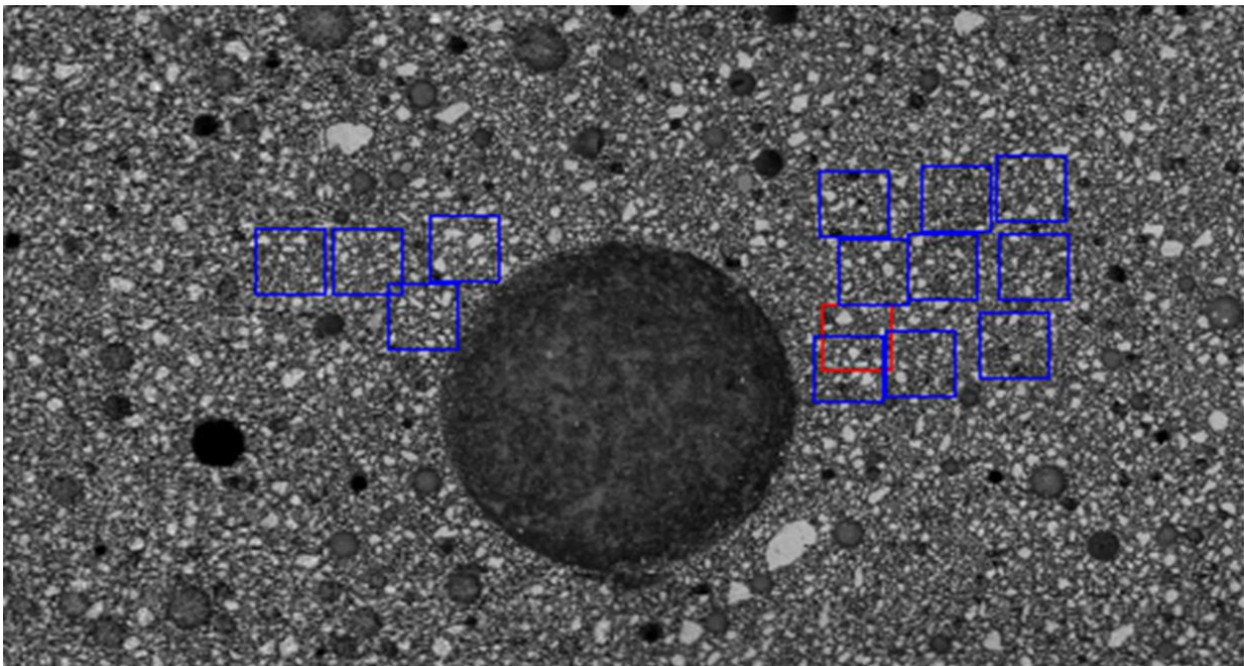


Figure 4-14: Fields Analyzed by UF and RJ Lee Group Inc.

The process of acquiring fields and processing them to create a segmented image is very labor intensive, has a fairly steep learning curve, and multiple fields must be collected to have statistically valid data for the sample. Work was performed in cooperation with RJ Lee Group Inc. to automate this process, with the purpose of reducing the time and operator skill required to obtain proper cement images.

A preliminary comparison study was conducted to investigate the potential for automated SEM field acquisition to reduce time and labor requirements for the process. The first part of this study involved the manual acquisition of 15 fields (BSE and EDS maps) of a polished cement

powder mount using the SEM equipment at the University of Florida. Upon completion of the analysis at the University of Florida, the specimen was then sent to RJ Lee group, and the same fields were located and acquired using their automated SEM equipment. The fields acquired are shown in Figure 4-14.

The acquisition of the BSE image and EDS maps at the University of Florida required approximately one hour of analyst time per field, with approximately 15 minutes of that hour being devoted to stage movement, refocusing, and BSE image acquisition, while the remainder of the hour was required to obtain the EDS maps themselves. This resulted in a total of roughly 15 hours of analyst time required for the acquisition of one set of fields for a single portland cement specimen.

The use of an automated SEM at RJ Lee Group allowed the identical fields to be located all at once, and placed in a queue to be analyzed. This meant that the analyst could step away after the fields to be analyzed were specified, and return with all of the images acquired. This required three hours of analyst time, which is a reduction of 80% (Watts, et al., 2013).

Table 4-4: Average Difference Between the Manual and Automated Image Sets

Phase	Area Fraction	Perimeter Fraction
C ₃ S	0.38%	0.81%
C ₂ S	2.75%	6.54%
C ₃ A	2.60%	6.17%
C ₄ AF	0.53%	1.20%

The data shown in Table 4-4 show the average difference between the data presented in Tables 4-2 and 4-3, and the data obtained from fields acquired at UF. The two data sets ended up agreeing fairly well, especially when the much higher resolution of the X-ray maps provided by RJ Lee Group was taken into account. This, combined with the inherently subjective nature of the segmentation process itself, means that the substantial reduction in acquisition time enabled by the use of an automated SEM does not degrade the accuracy of the data obtained (Watts, et al., 2013).

In order to ensure that the images were of equivalent quality and to prove that the acquisition of fields manually had no impact on the resulting image segmentation process, both

sets of images were processed in the same way at the University of Florida. The resulting difference between the two sets is described in Table 4-5.

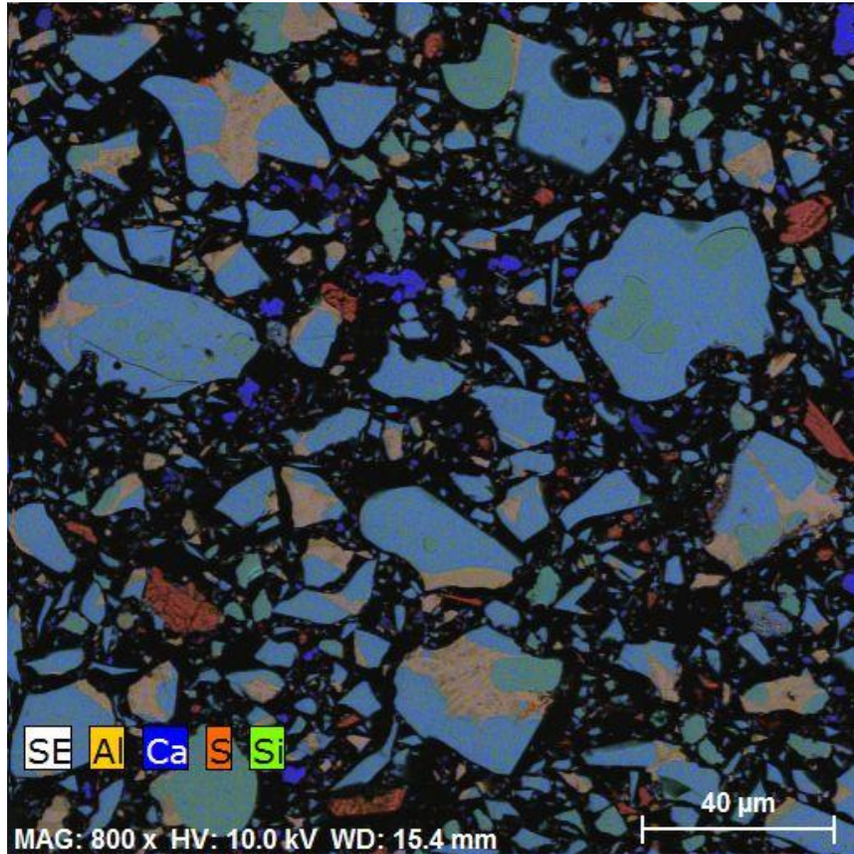


Figure 4-15: New False Color Image

After the comparison study, efforts continued at RJ Lee Group to further automate the characterization process. The ultimate goal of the collaboration was to completely automate the scanning and processing of cement samples. As of this research, some further improvements have been made to the acquisition process, but the image processing is still done manually. The chief improvement to the acquisition process was the creation of an image that is already colored in such a way that it allows all the major phases to be distinguished. An example image can be seen in Figure 4-15.

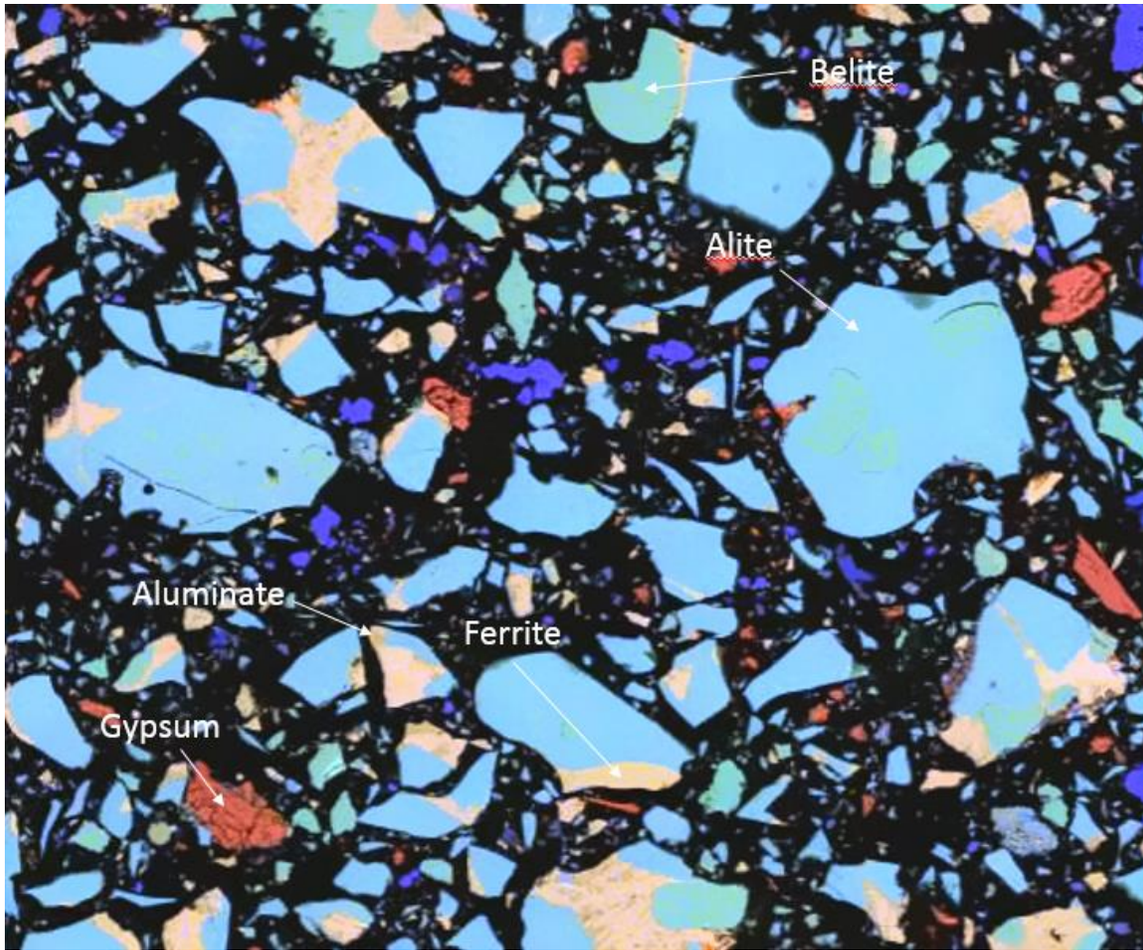


Figure 4-16: After Processing, Image Ready for Classification

This image was supplied directly from RJ Lee, and it eliminated the process of combining the x-ray maps and backscatter electron images. Different colors were assigned to different elements, as shown in the key at the lower left of Figure 4-15, and the resulting shades indicate the relative mass percentages of the elements present. The best example of this can be seen in the difference between alite and belite. Silicon was assigned to green, and calcium was assigned to blue, and because belite (C_2S) has a higher ratio of silicon to calcium, it showed up as a greener shade of blue-green, while alite appeared bluer.

This image was cropped, levels were adjusted, and the thresholded blur filter described earlier was applied to reduce noise. The resulting image, shown in Figure 4-16, was able to be classified in Multispec immediately. Image analysis proceeded as usual after this image was loaded into Multispec.

Several images were processed in this manner; however, instead of averaging the phase fractions as was done when the manual and automated fields were compared, the data from each field was looked at individually. The field from which the measurement most closely matched the XRD data was chosen to supply inputs to the VCCTL software. The phase volume fraction data measured from the images were compared to the data calculated from the XRD data obtained from the cement plant. Table 4-6 shows a comparison between the plant data and the data measured from the image shown in Figure 4-17.

Table 4-5: XRD Cement Phase Fraction vs. SEM Microanalysis Measurements

Phase	Mass % (XRD)	Volume % (XRD)	Volume % (Measured)	Surface Area % (Measured)
Alite	68.31	69.14	66.14	58.44
Belite	15.04	15.73	15.72	20.37
Aluminate	5.39	5.59	5.74	17.12
Ferrite	11.25	9.54	12.04	4.07

As Table 4-5 shows, the phase volume fraction data measured from Figure 4-17 is within 3% of the values obtained via XRD. These phase fraction data were used for all simulations of this cement.

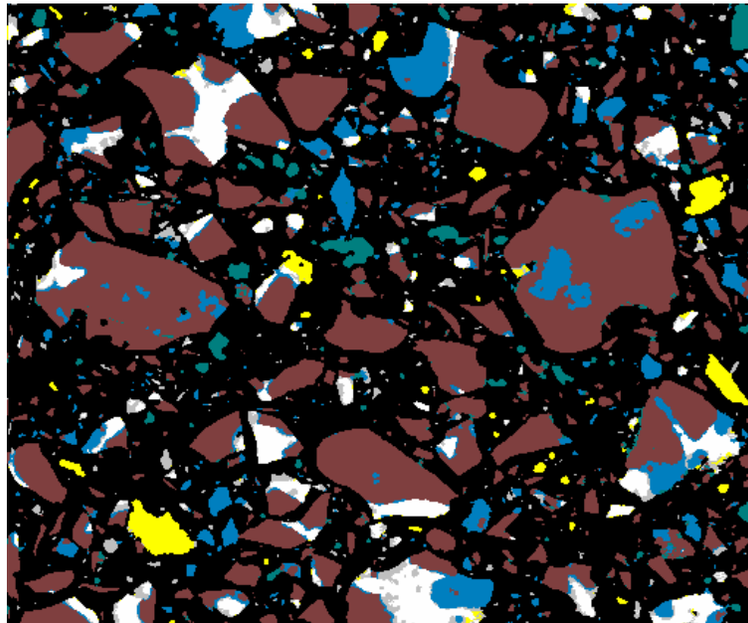


Figure 4-17: Final Cement Image, Brown Representing Alite, Blue-Belite, White-Ferrite, Grey-Aluminate, Teal-Free Lime, Black-Void, and Yellow-Gypsum

It was decided to utilize data from an individual field rather than the average of several fields because the availability of the mass fraction data from XRD (obtained in earlier analyses) revealed that the image analyses systematically overestimated the volume fractions of certain phases, particularly alite and belite. Since the accuracy of phase characterization is the ultimate goal for this technique, it was deemed acceptable to pick an individual field that matched the XRD data closely because of the standardized and well established nature of the XRD test.

CHAPTER 5 VCCTL SOFTWARE OUTPUT DATA

Overview

The outputs provided by the VCCTL software can be divided into two categories. There are properties that are computed and measured continuously as part of the hydration simulation, and are referred to as continuous measurements by the software. There are outputs that result from additional simulations calculated using a snapshot of the hydrated microstructure as an input, which are referred to as periodic measurements by the model because they can only be performed on static microstructures that have been saved at user specified intervals. The following list provides the potential outputs from the software:

- Cement and hydration product phase volume fractions;
- Solution concentration of sodium, potassium, calcium, sulfate;
- Solution activity of sodium, potassium, calcium, sulfate;
- Total porosity;
- Fraction porosity connected in the x, y, and z directions;
- Average fraction porosity connected;
- Fraction solids connected in the x, y, and z direction;
- Average fraction solids connected;
- Solution pH;
- Solution conductivity;
- Non-evaporable water content (ignited and unignited);
- Gel/space ratio;
- Binder temperature;
- Heat release;
- Degree of hydration;
- Cycles run;
- Elastic modulus of the paste;
- Elastic modulus of the concrete; and
- Strength of the concrete.

Of the potential simulated outputs available in the model, only average fraction solids connected, modulus of concrete, and strength of concrete are reported in this research.

Continuous Measurements

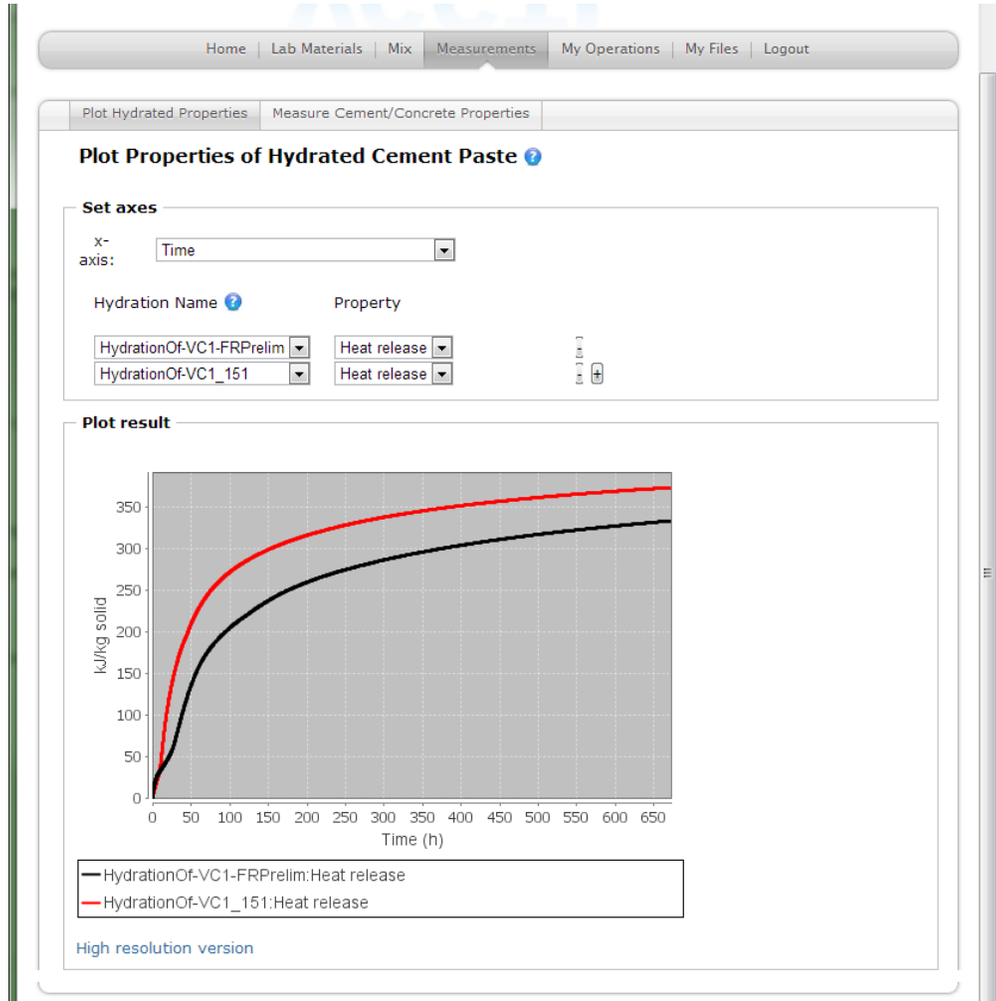


Figure 5-1: Continuous Measurements Display Page

The VCCTL software provides 64 continuous output parameters that describe the simulated hydrating cementitious microstructure. Of these 64 output parameters available, 34 describe the volume fractions of the cement and hydration products that make up the physical composition of the microstructure. The other 30 outputs include solution concentrations and activities for sodium, potassium, calcium and sulfate, total porosity, fraction porosity three-dimensional connectivity of solids, average fraction porosity, pH and conductivity of the pore solution, non-evaporable water content, gel/space ratio, temperature of the binder, heat release,

degrees of hydration for cement and fly ash, and the number of simulation cycles run. These outputs provide a wealth of fundamental information about the hydrating cement paste. For example, Figure 5-1 displays the resultant heat release of a typical simulated microstructure during hydration.

Some of these outputs, specifically those that simulate pH, conductivity, and ionic composition of the pore solution, are intended to provide information that can be linked to the durability of concrete via degradation modeling (Bullard, 2002). Some work has been performed utilizing ion chromatography of extracted pore solution to evaluate the accuracy of the VCCTL with regard to prediction of ionic concentrations (Feng, 2004). The software itself, however, provides no direct provision for the prediction of concrete durability or service life.

Some of the outputs provided by the model have been used for the experimental verification of the accuracy of the hydration model. The volume fraction predictions, for instance, have been compared to experimental values obtained via in situ x-ray diffraction techniques (Valentini et al, 2013) and prediction of degree of hydration has been verified experimentally (Bentz, 2006 and Bentz, et al., 2000) via measurement of non-evaporable water within the system. The quantity of heat release as simulated by the model has been compared to experimental measurements of heat of hydration via ASTM C186, and been used successfully for verification of the model (Bentz, et al., 2000).

Some of the measurements provided as continuous outputs have seen direct applications for prediction of concrete performance. The most important of these is the fraction solids connected value because it describes the percentage of the solid microstructure that has percolated through the simulated volume over time. Fraction of solids connected can be used to describe the phenomena of 'set', that is, the linking up of individual hydrating cement grains to form a rigid, interconnected matrix.

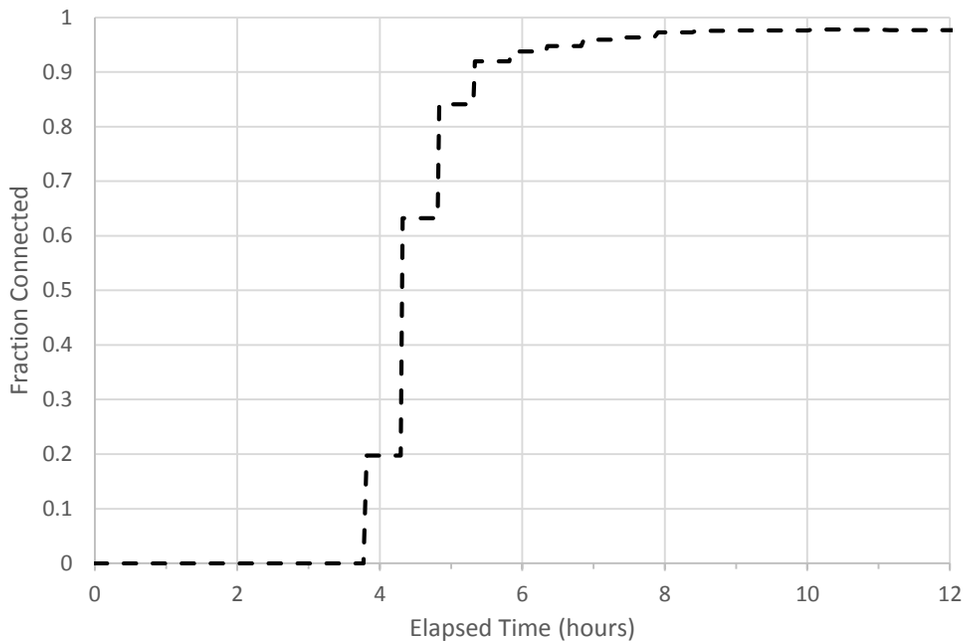


Figure 5-2: Fraction Solids Connected Curve

Experimental tests for the setting time of cement typically adopt arbitrary limits to define “initial set” and “final set.” As described in ASTM C403, time of set is determined by the resistance of a mortar to the penetration of a needle with a specific cross sectional area, and defines initial set as a resistance of 500 pounds per square inch, and final set as 4000 pounds per square inch. The values for defining initial set and final set for the fraction solids connected value can be assigned in a similarly arbitrary fashion. This method has been used in industry applications to evaluate the effects of temperature on set time (Bullard, et al., 2013) with some success. For this research, the value for initial set was defined at a fraction solids connected value of 0.2, and final set was defined at 0.8. An example of the fraction solids connected output from the VCCTL software can be seen in Figure 5-2.

Research has been performed to verify the prediction of the compressive strength of mortar cubes at later ages via the gel/space output provided by the model and the Power’s gel/space theory (Garboczi, et al., 2004). This method is somewhat limited, as it requires experimental measurements at early age to calibrate the empirical theory and enable later-age predictions. It is also noteworthy that this is not a built-in function of the VCCTL software, but an application of one of its outputs.

Periodic Measurements

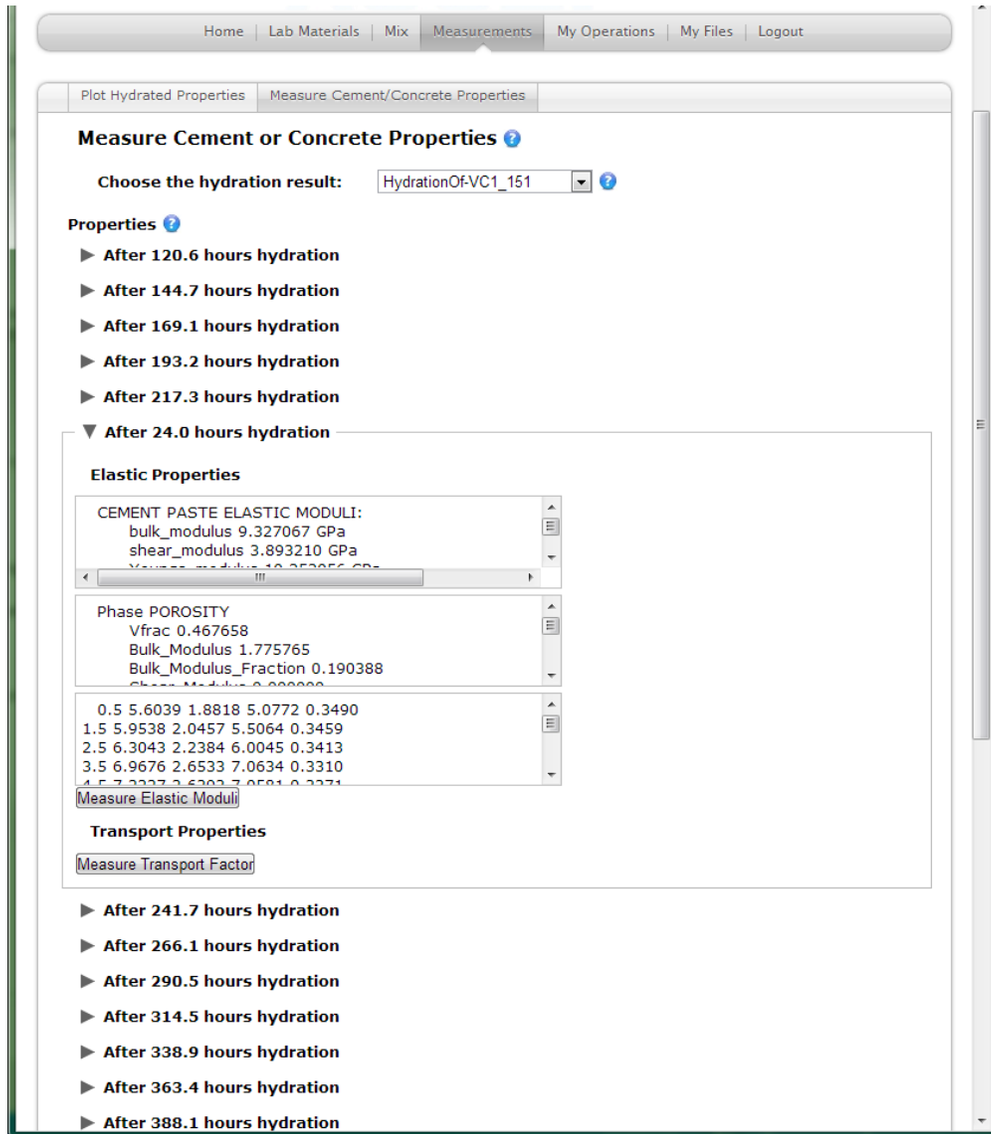


Figure 5-3: Periodic Measurement Display Page

The second category of measurements are those obtained from calculations run on hydrated microstructures of various ages. It is from these outputs that elastic modulus and strength properties are obtained. The measurements screen for periodic outputs is shown in Figure 5-3.

Elastic Modulus

The VCCTL software computes the elastic moduli of hydrated cement paste, mortar or concrete and is first computed from the hydrated microstructure using a finite element model.

The resulting modulus of the paste is then used as an input for another, slightly different finite element calculation which incorporates the modulus of the concrete, based on the computed paste modulus as well as the input bulk and shear moduli for the aggregates used. A summary of the methods by which these computations are executed is summarized in the following sections.

Hydrated Cement Microstructure Modulus

The elastic modulus of hydrated cement paste is computed using an effective medium theory (EMT) for three-dimensional spherical particles (Garboczi & Day 1995). This theory calculates the bulk and shear moduli of the composite paste using known values of the bulk and shear moduli and volume fractions of each specific phase present in the paste. The equations used to calculate the effective shear and bulk moduli of the paste are as follows:

$$\frac{1}{G} = \frac{c_1}{G+\beta(G_1-G)} + \frac{c_2}{G+\beta(G_2-G)} \cdots + \frac{c_n}{G+\beta(G_n-G)} \quad (5-1)$$

$$\frac{1}{K} = \frac{c_1}{K+\alpha(K_1-K)} + \frac{c_2}{K+\alpha(K_2-K)} \cdots + \frac{c_n}{K+\alpha(K_n-K)} \quad (5-2)$$

$$\beta = \frac{2(4-5\nu)}{15(1-\nu)} \quad (5-3)$$

$$\alpha = \frac{2(4-5\nu)}{15(1-\nu)} \quad (5-4)$$

Where:

- G and G_n , K and K_n are the shear and bulk moduli (GPa) of the composite and n^{th} phase;
- ν is the Poisson's ratio of the composite;
- c_n is the volume fraction of the n^{th} phase; and
- α and β are dimensionless constants.

An algorithm which utilizes equations (5-1) - (5-4) simulates the composite elastic modulus of the three-dimensional microstructure by treating the digital image as a mesh of linear finite elements, where each voxel is an element. Individual phase properties are assigned to each voxel, and composite properties are computed by applying a strain and computing the appropriate energy averages (Garboczi & Day, 1995).

Concrete and Mortar Modulus

The resulting modulus of the paste is then used to calculate the modulus of the concrete. This calculation uses a more complex differential EMT, which models the concrete as a matrix with spherical inclusions. The inclusions are composite elements composed of individual aggregate particles that are surrounded by thin shells of altered matrix material (representing the interfacial transition zone). The aggregates and surrounding shells are mapped as a single particle with a diameter equal to that of the outer shell. The effective or composite moduli are calculated from the moduli of the particle and shell. The equations to perform this mapping are quite lengthy, and detailed in Appendix A.

The new effective particles are larger than the original aggregate particles, and thus the original particle volume fractions must be normalized to account for the change in diameter. The equations used to perform this normalization are as follows:

$$\alpha_j = \frac{a_j^3}{b_j^3} \quad (5-5)$$

$$c' = c \sum_{j=1}^M f_j \alpha_j \quad (5-6)$$

$$f'_j = \frac{f_j \alpha_j}{\sum_{j=1}^M f_j \alpha_j} \quad (5-7)$$

Where:

c = original volume fraction;

c' = normalized volume fraction;

f_j = original partial volume fraction of the j^{th} type of particle lying in a certain diameter range;

f'_j = normalized partial volume fractions of the j^{th} type of particle lying in a certain diameter range;

b_j = the diameter (in microns) of the original particle; and

a_j = the diameter of the j^{th} type new effective particle.

The resulting bulk and shear moduli are solved for numerically using a set of differential equations detailed in Appendix A. This numerical method was verified using the same method by which the cement paste modulus is calculated, that is a three-dimensional digital image was built with the aggregates and their surrounding zones of altered matrix material, and the effective

modulus of the image was found using finite element simulation. Excellent agreement was found between the two methods. (Garboczi & Berryman, 2001).

Compressive Strength

The VCCTL software finds the compressive strength of concrete via a direct, empirically-derived relationship. The available literature on this relationship points to the ACI 318 (American Concrete Institute, 2010) code as the origin of this equation (Garboczi, et al., 2004). From ACI 318, an equation for the calculation of compressive strength can be found. When used with metric units, this equation takes the following form:

$$E_c = 4.73\sqrt{f'_c} , \quad (5-8)$$

where f'_c and E_c are compressive strength in MPa and elastic modulus in GPa, respectively.

Solving this equation for compressive strength yields

$$f'_c = \left(\frac{E_c}{4.73}\right)^2 . \quad (5-9)$$

Or if the equation is rearranged

$$f'_c = 0.0447E_c^2 . \quad (5-10)$$

The software source code however, shows a different relationship between strength and modulus, which takes the following form:

$$f'_c = 0.0006E_c^{2.9664} . \quad (5-11)$$

Figures 5-4 through 5-6 show example strength and modulus data from simulations. In Figure 5-6, it can be seen that the relationship between strength and modulus is indeed that of the equation from the source code. The commentary relating to this equation within the source code also indicates that the strength values provided from this empirical relationship are for large cubes of unspecified size of the type commonly used in Europe. Literature on the relationship between concrete cylinder and cube strength (Elwell & Fu, 1995) indicates that the strength of cylinders is approximately 0.8 that of cubes; however, for this study the output of the model was compared directly to measured values as discussed in Chapter 8.

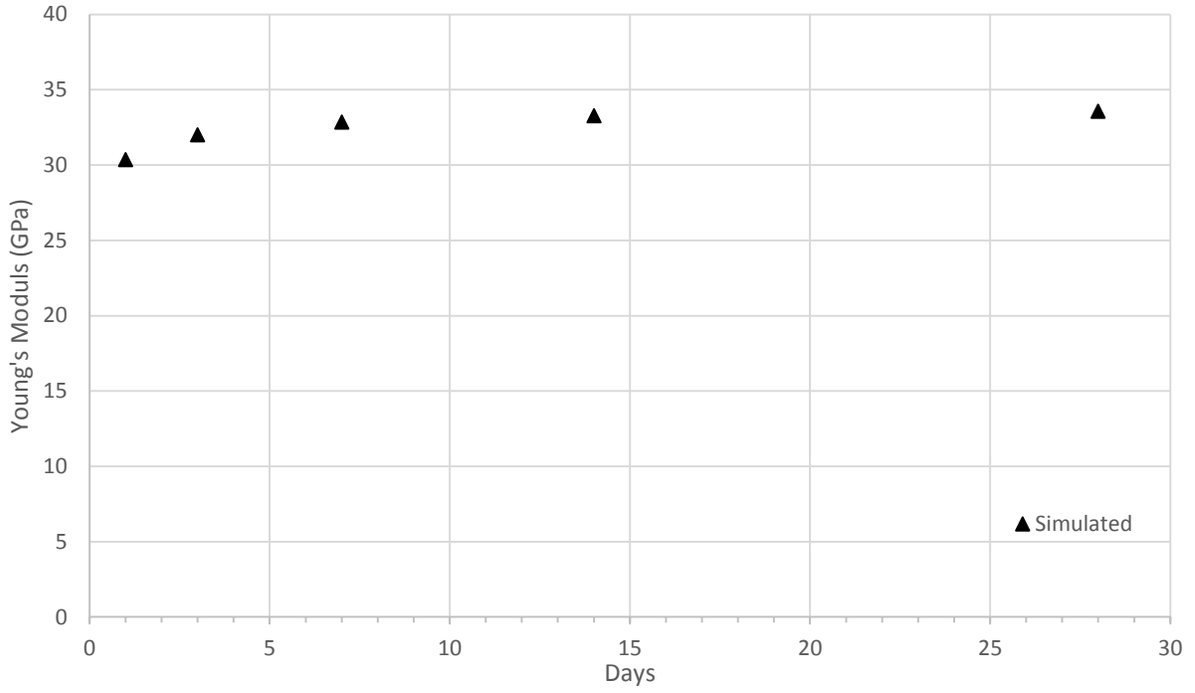


Figure 5-4: Predicted Young's Modulus vs. Time, w/c 0.4

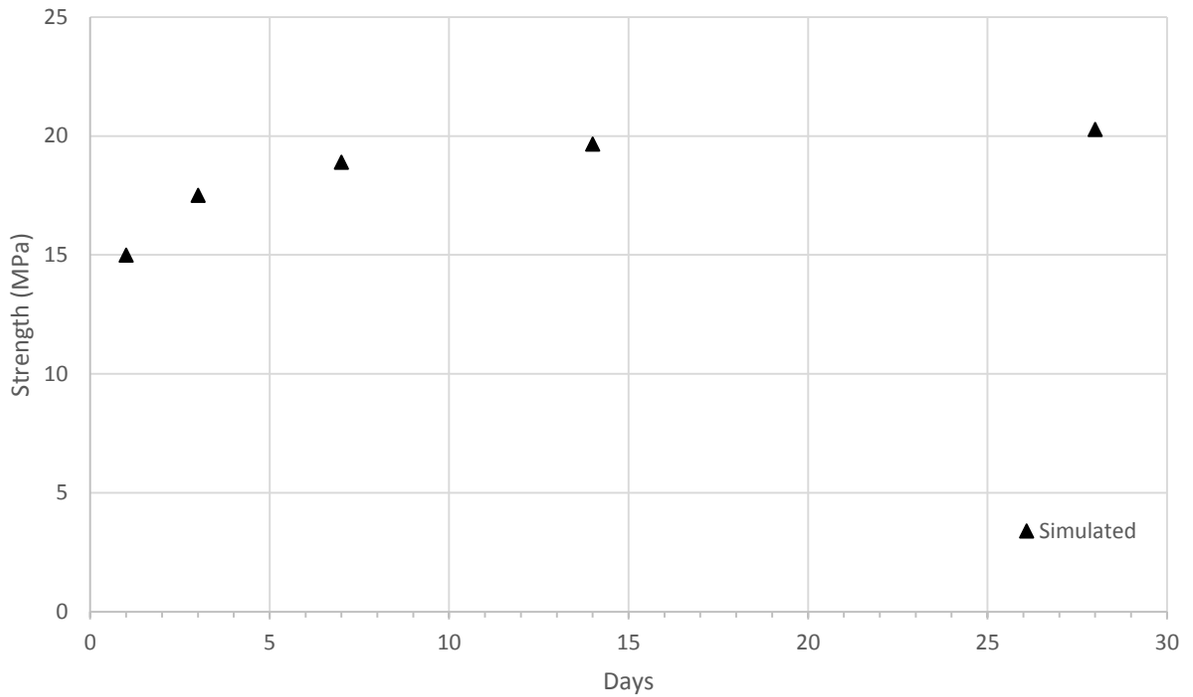


Figure 5-5: Predicted Strength vs. Time, w/c 0.4

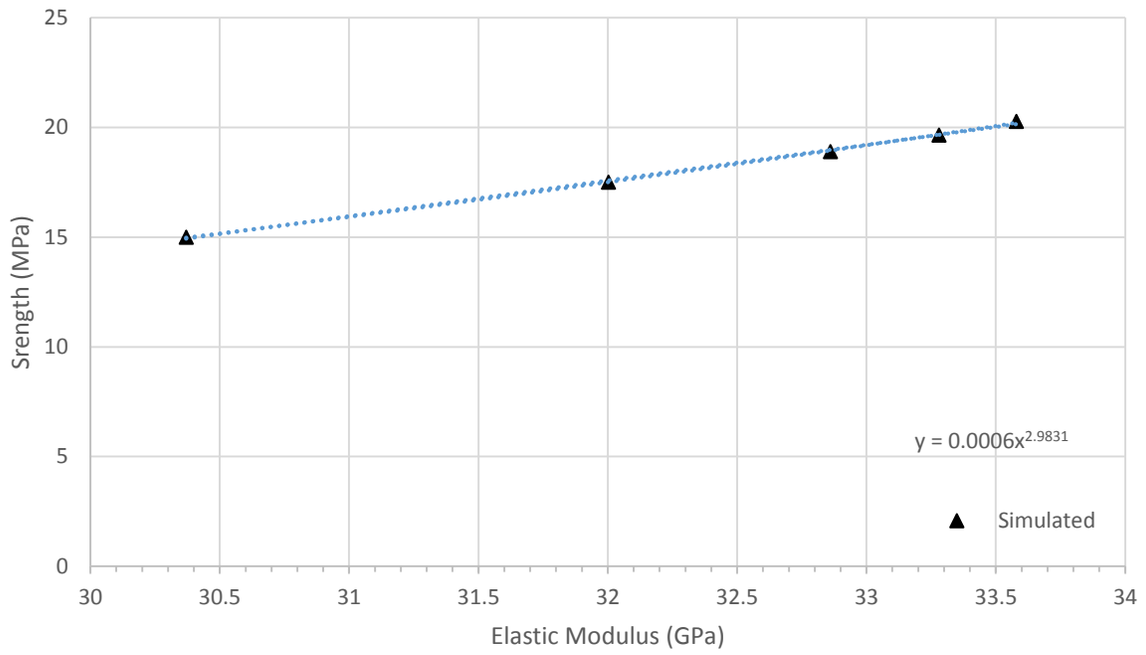


Figure 5-6: Strength vs. Elastic Modulus, w/c 0.4

CHAPTER 6 PHYSICAL TEST PROGRAM

Physical Testing Program

All experiments relied on physical data from real concrete to evaluate the effectiveness of the model. A summary of the different mixes used in this research can be seen in Table 6-1. The densities for the aggregates shown were in saturated, surface dry (SSD) condition.

Table 6-1: Mixture Design Summary

Material	Mix 1 (0.4 w/c)	Mix 2 (0.45 w/c)	Mix 3 (0.5 w/c)	Mix 4 (0.55 w/c)	Mix 5 (0.4 w/c)	Mix 6 (0.4 w/c)	Mix 7 (0.4 w/c)
Cement (lb/yd ³)	681	681	681	681	681	681	681
Water (lb/yd ³)	272	306	341	375	272	272	272
Fine Agg. (lb/yd ³)	1093	1061	1030	995	1093	1093	1093
Coarse Agg. (lb/yd ³)	1681	1631	1586	1532	1681	1681	1681
Admixture	-	-	-	-	Type D	Type F	Type F
Dosage (oz/cwt)	-	-	-	-	4.0	2.0	4.0

The cement and aggregates used in each of the mixes were obtained from a single source to ensure consistency. Since the primary focus of the research was to determine the influence of the different inputs on simulation accuracy, a range of mixes with different water contents was made. These mixes had the raw material additions listed in Table 6-1, with water-to-cement ratios of 0.4, 0.45, 0.5 and 0.55. The difference in the volume of water for each mix was accounted for by changing the total aggregate content while keeping the ratio of coarse-to-fine aggregate the same.

Physical specimens were also created with different dosages of two commonly used water-reducing admixtures, a Type D mid-range water reducer and retarder, and a Type F high-range water reducer. Mix 1, with a w/c of 0.4, was the control mix and admixtures were included to create Mixes 5, 6 and 7.

Specimens were prepared following the standard procedures outlined in ASTM C192. Prior to batching, an approximate amount of coarse aggregate was bagged and soaked for at least 24 hours. The bags were removed from the soak an hour prior to batching to allow excess water to drain. While the coarse aggregate was draining, the cement and fine aggregate were weighed and put into buckets. The coarse aggregate was then batched, with a small amount of aggregate

set aside from each bag for determination of moisture content. The aggregate set aside was weighed wet, then put into the oven overnight to remove all moisture following the procedures outlined in ASTM C566. The following day the water content and coarse aggregate contents were adjusted based on the difference between the measured moisture content and the SSD moisture content.

A small initial concrete batch, approximately 10% the size of the primary batch, was prepared; this served to coat the inside of the mixer with mortar, and prevent any mortar loss of the primary mix. Coarse and fine aggregates along with approximately half the water were then added to the mixer and mixed until evenly distributed. The cement and remaining water were added and mixed for three minutes. The concrete was then allowed to rest for approximately three minutes. Following this step the admixture (if any) was added and the concrete was mixed for another three minutes.

Each of the concrete batches were tested for unit weight (ASTM C138), temperature (ASTM C1064), and air content (ASTM C138) in accordance with the ASTM standard method for each respective test. Mixes containing admixtures also had a portion of their mortar sieved out and placed into molds for the time of set test (ASTM C403). The results for the air content, unit weight, temperature, and slump tests are shown in Table 6-2. Mixes three and four were too fluid for the slump test to be performed to ASTM standards.

Table 6-2: Summary of Test Results for Fresh Concrete

Material	Mix 1	Mix 2	Mix 3	Mix 4	Mix 5	Mix 6	Mix 7
Temperature (°F)	75	72	74	75	76	74	74
Air Content (%)	2.5	2.25	0.3	0.0	3.0	3.0	3.75
Unit Weight (lb/yd ³)	143.4	142.2	139.1	137.7	141.3	140.5	142.16



Figure 6-1: Preparation of Cylinders

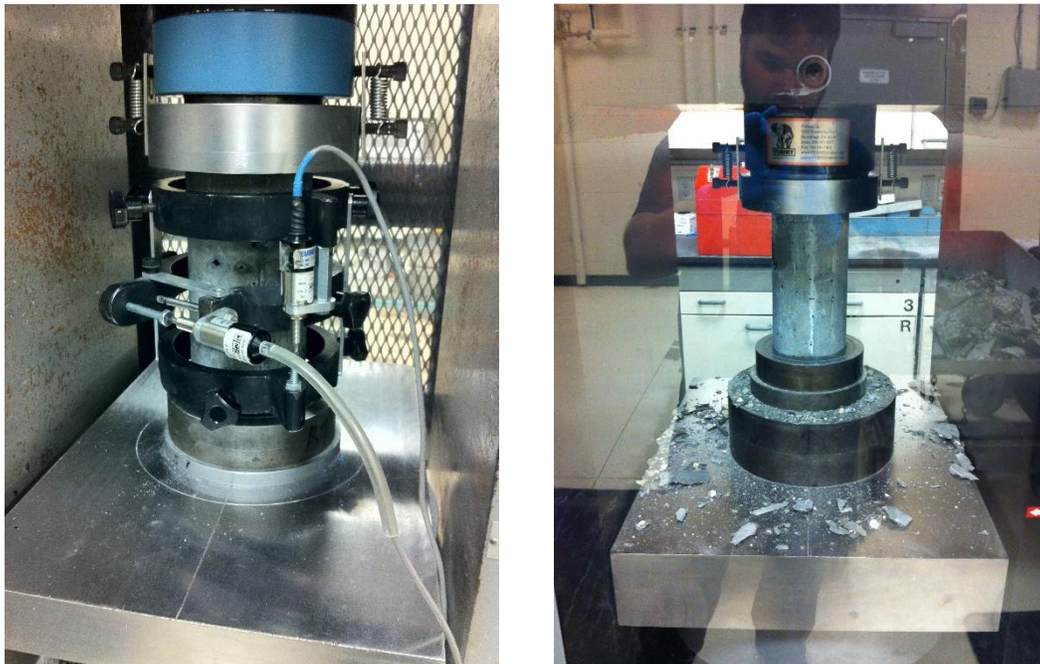


Figure 6-2: Elastic Modulus and Compressive Strength Testing

Each batch of thirty-eight 4" by 8" test cylinders were used for the creation of specimens for the testing of compressive strength and modulus in accordance with ASTM C192. Cylinders

were filled in two lifts, with a vibratory table used for 15 seconds between each lift to consolidate the concrete as depicted in Figure 6-1. Cylinders were then capped and cured for 24 hours prior to stripping the molds and the first strength and modulus tests. The time of set test was run in accordance with ASTM C403, penetration testing began at two hours, after which penetration resistance measurements were taken at regular intervals. The apparatus for the penetration resistance test is shown in Figure 6-3.

Hardened cylinders were ground flat on both ends and tested for compressive strength and elastic modulus in accordance with ASTM C39 and ASTM C 469, respectively. Figure 6-2 provides an illustration of these tests. Strength and modulus values were obtained at 1, 3, 7, 14, and 28 days in order to provide a wide range of time scales for the evaluation of the accuracy of the VCCTL software.



Figure 6-3: Time of Set Test Apparatus

Isothermal Calorimetry

Isothermal calorimetry data were collected for each of the mixes that contained admixtures, as well as the control mix with the w/c of 0.4. For each cement, water, and

admixture batch, three samples were mixed and measured. Thermal masses of cement and water for each sample were calculated such that the total heat capacity of the mixture was equal to that of the reference sample. Admixtures and water were proportioned by mass to reflect the actual amounts present in the concrete samples. The mixture was then loaded into the admix ampoule syringes by mass and were allowed to equilibrate overnight prior to mixing, after which data was collected for 168 hours (7 days). A summary of tests and ages at which they were performed for all mixes is shown in Table 6-3.

Table 6-3: Tests and Ages (Days)

Test	Mix 1 (0.4 w/c)	Mix 2 (0.45 w/c)	Mix 3 (0.5 w/c)	Mix 4 (0.55 w/c)	Mix 5 (0.4 w/c)	Mix 6 (0.4 w/c)	Mix 7 (0.4 w/c)
ASTM C39 Compressive Strength	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28
ASTM C469 Elastic Modulus	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28	1, 3, 7, 14, 28
ASTM C403 Time of Set	1	-	-	-	1	1	1
ASTM C1702 Heat of Hydration	1-7	-	1-7	1-7	1-7	1-7	1-7

CHAPTER 7 COMPARISON OF PHYSICAL TEST DATA TO MODEL OUTPUTS

Experimental Design

The primary objective of the experiments conducted for this research was to validate the accuracy of the VCCTL software with regard to the simulation of the structural properties of concrete (as opposed to mortar or grout), and assess its potential value as a tool for the development of new concrete mixture designs. The vast majority of concrete mixes made today contain both supplementary cementitious materials (SCMs), such as fly ash and silica fume, and practically all mixes contain some type of chemical admixture. While mixes containing these materials can be simulated by the software, the characterization of these materials requires the development and refinement of procedures at least as complex as those for portland cement. The scope of this project precluded the use of SCMs for the validation study, however a method by which certain effects of some admixtures may be simulated was developed and investigated.

Currently, the VCCTL software does not possess the capability to properly model concrete mixtures which incorporate supplementary cementitious materials; therefore, the concrete mixture designs considered for the validation study contained portland cement as the only cementitious component. With this restriction imposed, the decision to test the accuracy of the simulations at a range of water-to-cement ratios was made. Water-to-cement ratio is widely known (Mindess & Young, 1981) to be the most important variable that affects the strength and elastic modulus of concrete. Concrete mixtures incorporating water-to-cement (w/c) ratios of 0.4, 0.45, 0.5, and 0.55 were selected, with the lower bound of 0.4 being the limit of feasible mixing without any water-reducing admixture, and the upper limit of 0.55 being above the designed water content of most mixes used in industry applications (Florida Department of Transportation, 2014). The experimental results for concrete mixtures created in the laboratory were compared directly to the compressive strength and compressive modulus elasticity data predicted by the software.

It was determined that investigating the influence of each of the different material inputs accepted by the VCCTL software would be important for evaluating its usefulness as a concrete mix design development tool. A control mix with a water-to-cement ratio of 0.4 was modeled first using all available input data, then several simulations were performed on the same mixture with a different non-essential input removed in each simulation. Because compressive strength is calculated by the software from the modulus, based on a fixed empirical relationship, it was not

considered when comparing the influence of different input data. For the removal of inputs which influence the hydration simulation, changes to the fraction solids connected curve were also investigated.

Experiments were conducted to investigate the accuracy of simulation of some concrete mixtures containing water-reducing admixtures which can increase the set time of fresh concrete as discussed in Chapter 3. The effect can be measured in fresh concrete using the time of set test, and can also be quantified using isothermal calorimetry. Since the VCCTL software utilizes heat of hydration data to calibrate the time scale of a simulation, it was decided to investigate the difference in set time obtained when isothermal calorimetry data were used from the hydration of a cement with a water-reducing admixture. Additionally, it was postulated that the long-term effects of the addition of a water-reducing admixture on strength and elastic modulus were negligible when mixes of the same water-to-cement ratio were compared (Mamlouk & Zaniewski, 2010). Given that this postulate holds true, this indirect method of accounting for admixture behavior could prove to be a substantial enhancement to the practicality of the software since the fraction solids connected value has been the primary output of the model used to predict the changes in set time.

Simulation Procedures

Except for the mixes evaluating the sensitivity of the VCCTL software to the absence of non-essential inputs as defined in Chapter 4, the mixes were simulated with the available, measurable inputs. These inputs included cement phase volume and surface area fraction data obtained via the SEM microanalysis method, the mass fraction data obtained from XRD analysis for the different phases of gypsum, particle size distribution curves, isothermal calorimetry curves, aggregate bulk and shear moduli, and aggregate shape data from the library of built-in materials. Simulations were also performed with real particle shape data for the cement, which was selected from the material library within the software database, based on its resemblance to segmented imagery.

To simulate the effects of water-to-cement ratio, the only changes made to each mix were the calorimetry curves, and the relative proportions of the mixes. Calorimetry data for the mix with 0.45 water-to-cement ratio were unavailable; the calorimetry data used for the mix at 0.4 water-to-cement were used as a substitute. The aggregate mass fractions, cement content, and water-to-cement ratio were modeled exactly as mixed. To gauge the sensitivity of the simulations

to the different non-essential inputs, several simulations were performed with selected inputs absent or disabled. The simulations used to evaluate input sensitivity were based on a control mix with a 0.4 w/c ratio. These included a mix with no calorimetry data, a mix with no real particle shape data, a mix with no aggregate shape data, as well as a mix with a microstructure 50% larger (on a side) than normal. The influence of aggregate moduli was also investigated by running three different Young's moduli in three individual simulations.

For the investigation of set time simulation using measured isothermal calorimetry data, four mixes were simulated with calorimetry data from three different admixture dosage combinations. The effects of a Type D (mid-range) water reducer and retarder, at a dosage of 4 ounces per hundred pounds of cement (4 oz/cwt); Type F high-range water reducer, at dosages of 2 and 4 oz/cwt; and a control mixture, with no admixture present. Isothermal calorimetry curves from each of these combinations were used in the simulations, and the mixes with admixtures present had cement particle dispersion enabled. The physical test results of the four different w/c are shown in Figures 7-1 and 7-2.

Results

Water-to-Cement Ratio

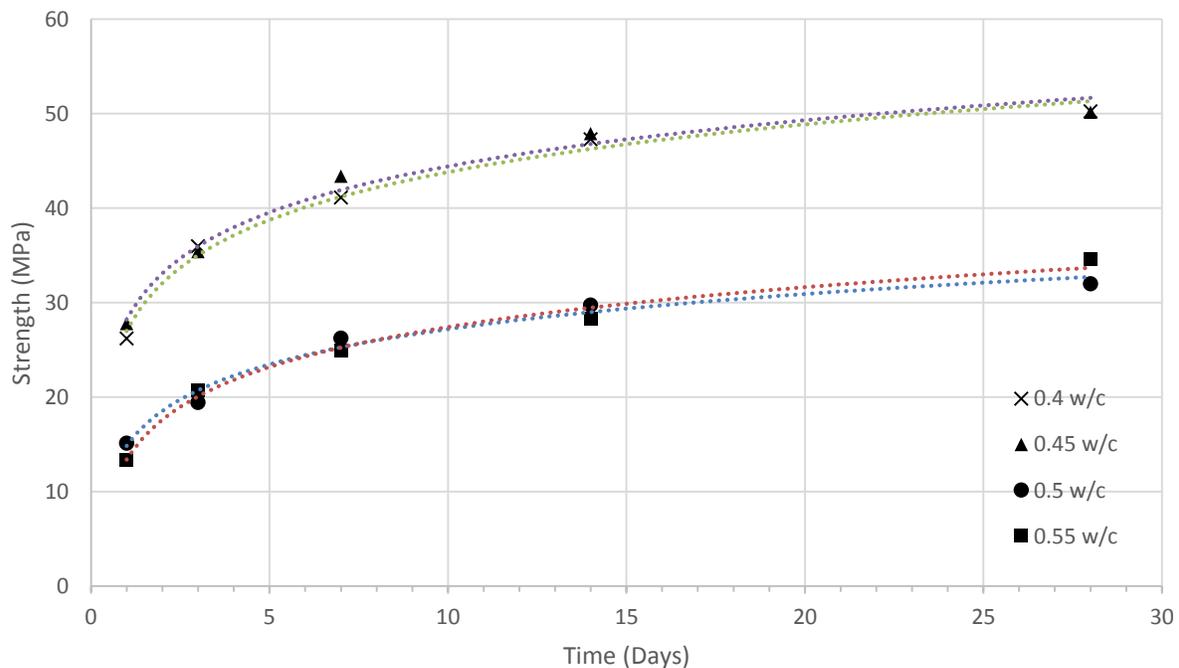


Figure 7-1: Compressive Strength vs. Time for Different w/c Ratios

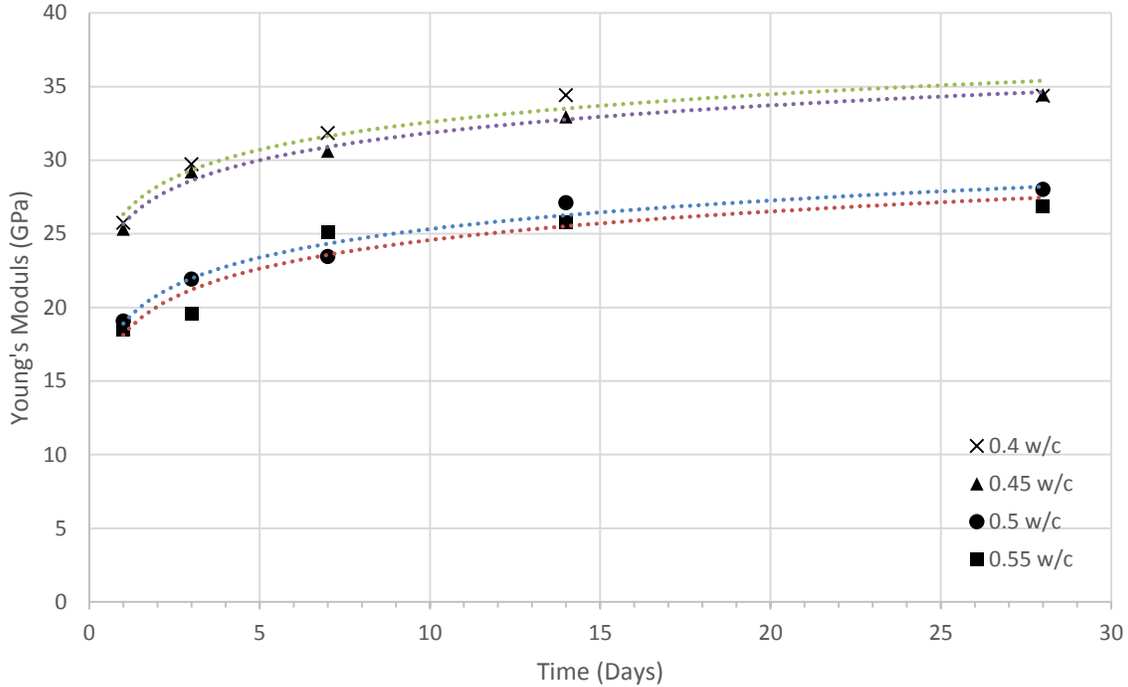


Figure 7-2: Young's Modulus vs. Time for Different w/c Ratios

Water-to-cement ratio (w/c) is one of the primary factors which affect compressive strength, modulus of elasticity, and other properties of the microstructure, which affect the general quality of concrete (Neville, 2011). The results presented in Figures 7-1 and 7-2 indicate that the measured strength and elastic modulus of the mixes increased as the water-to-cement ratio decreased.

Power and heat of hydration curves obtained via isothermal calorimetry for the different w/c are shown in Figures 7-3 and 7-4. Figures 7-3 and 7-4 indicate that while there was very little apparent difference between the maximum power outputs at the three different w/c, the ultimate energy reached at 7 days (7-day heat of hydration) increased slightly with an increase in w/c.

Figures 7-5 through 7-12 show comparisons of the strength and modulus results obtained from physical testing to the simulated results calculated by the model. The error bars for the physical test data represent plus or minus one standard deviation of the sample set average for each measurement.

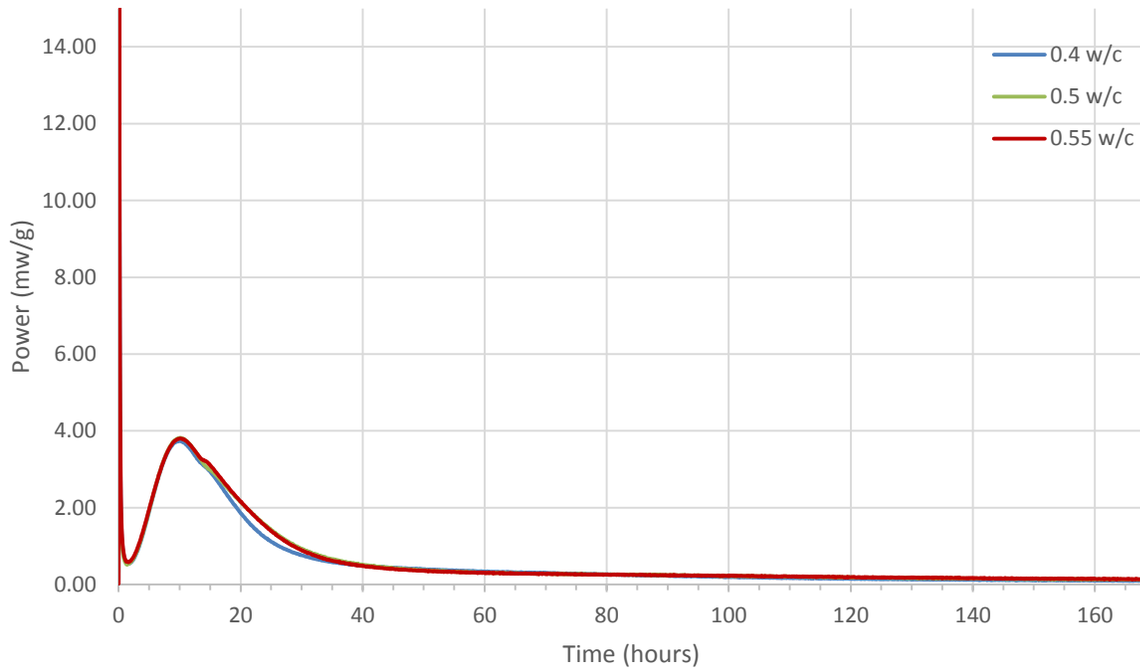


Figure 7-3: Power vs. Time for Different w/c

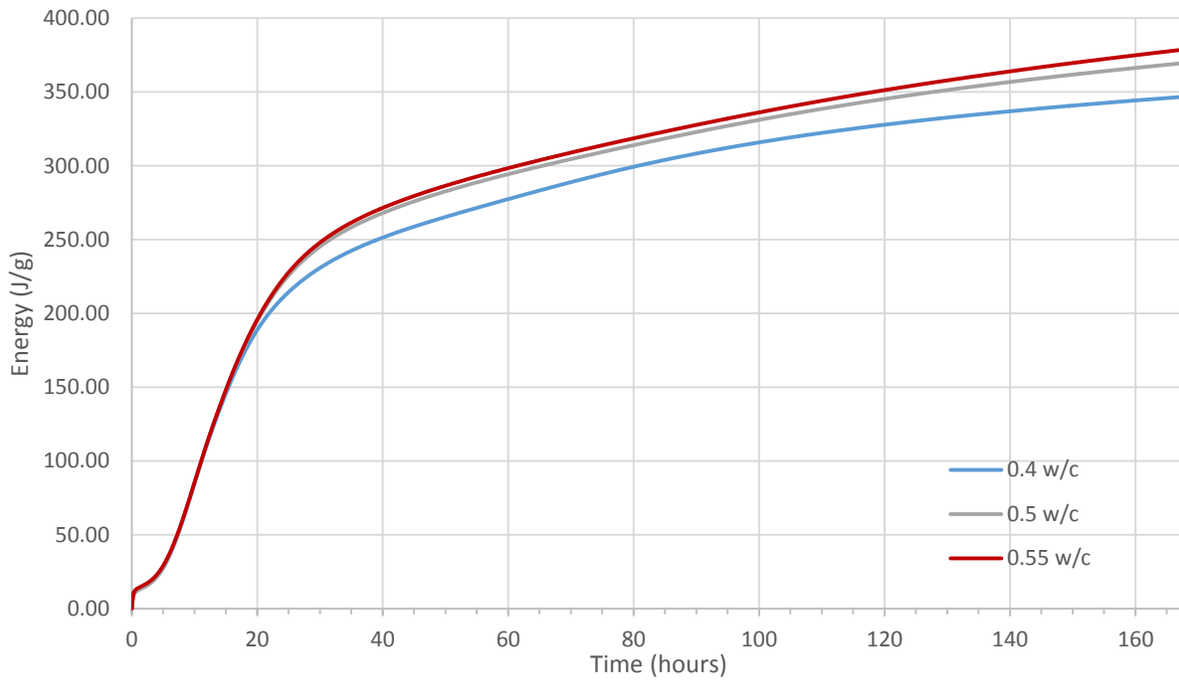


Figure 7-4: Energy vs. Time for Different w/c

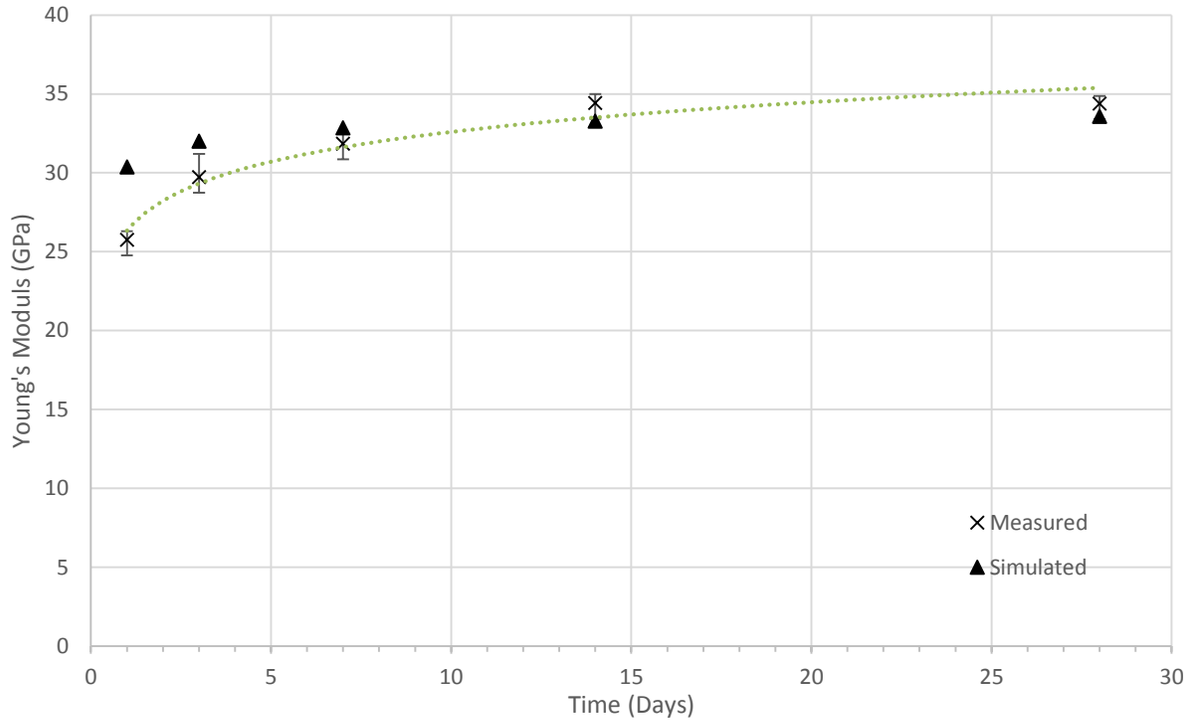


Figure 7-5: Young Modulus vs. Time, w/c of 0.4

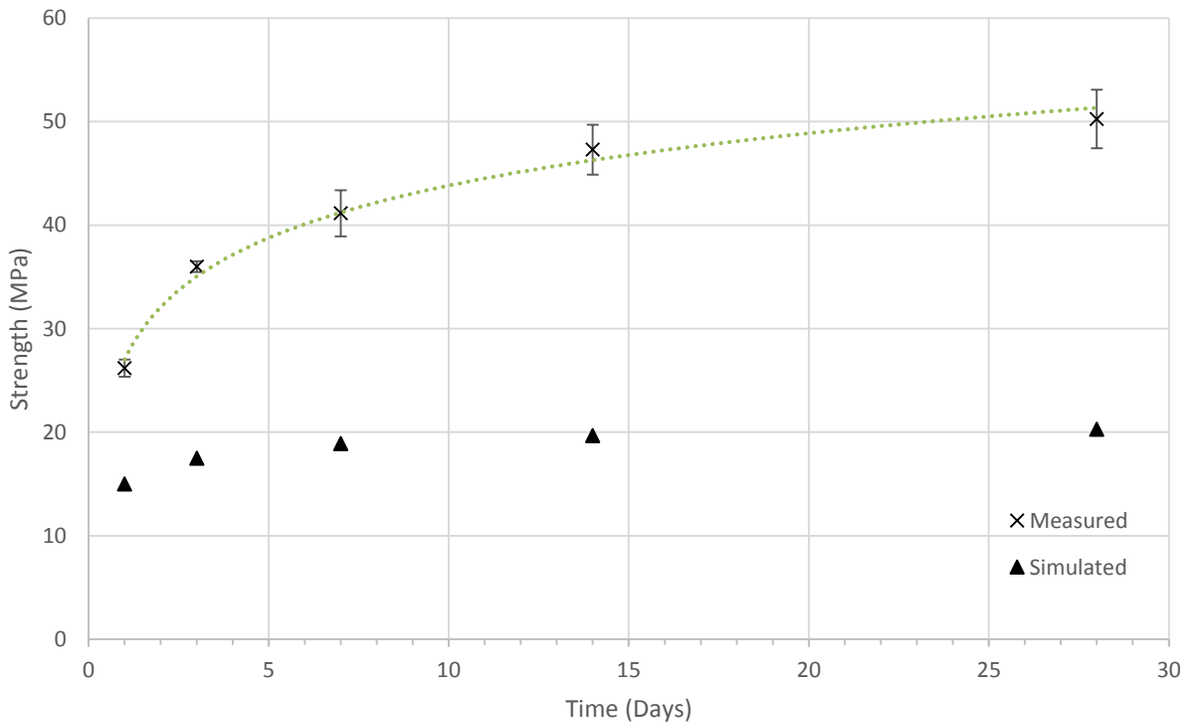


Figure 7-6: Compressive Strength vs. Time, w/c of 0.4

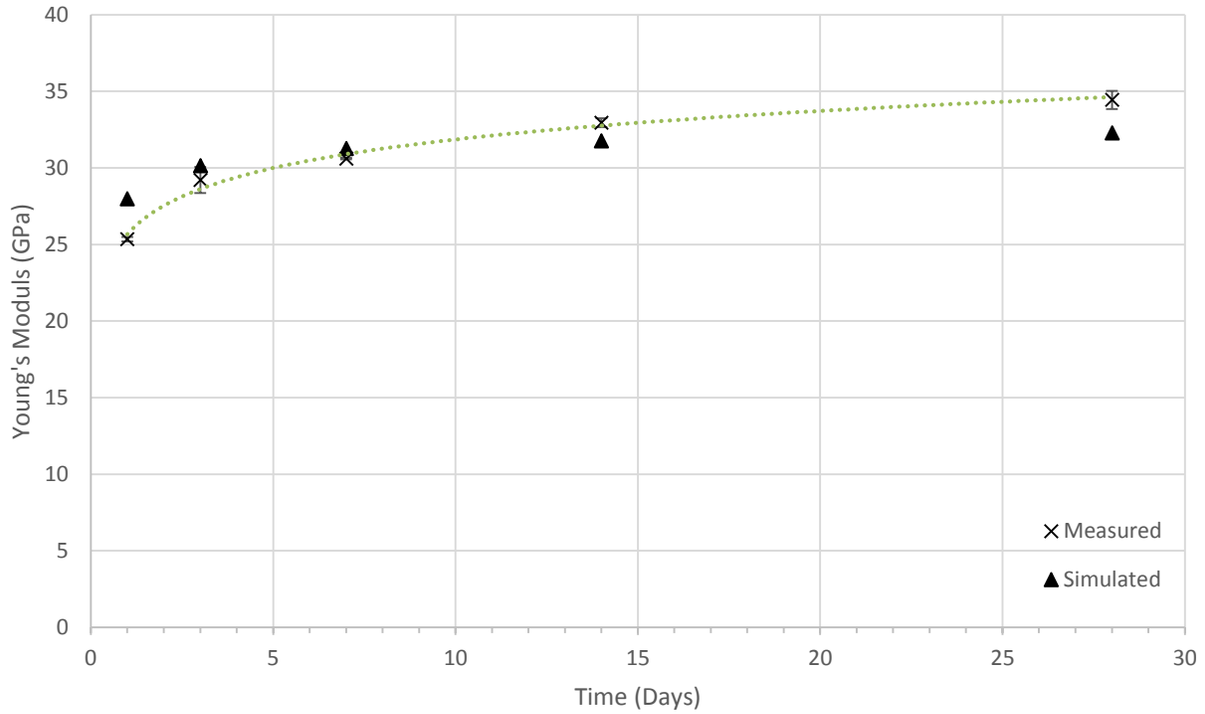


Figure 7-7: Young Modulus vs. Time, w/c of 0.45

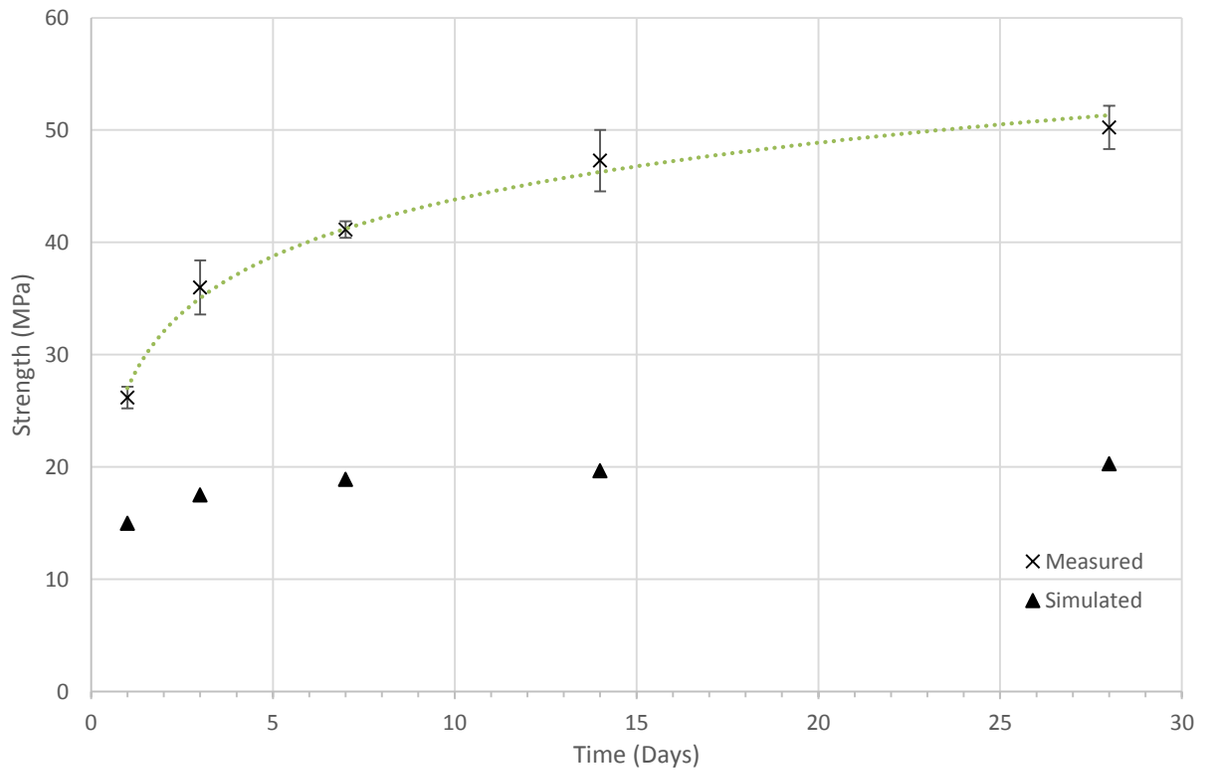


Figure 7-8: Compressive Strength vs. Time, w/c of 0.45

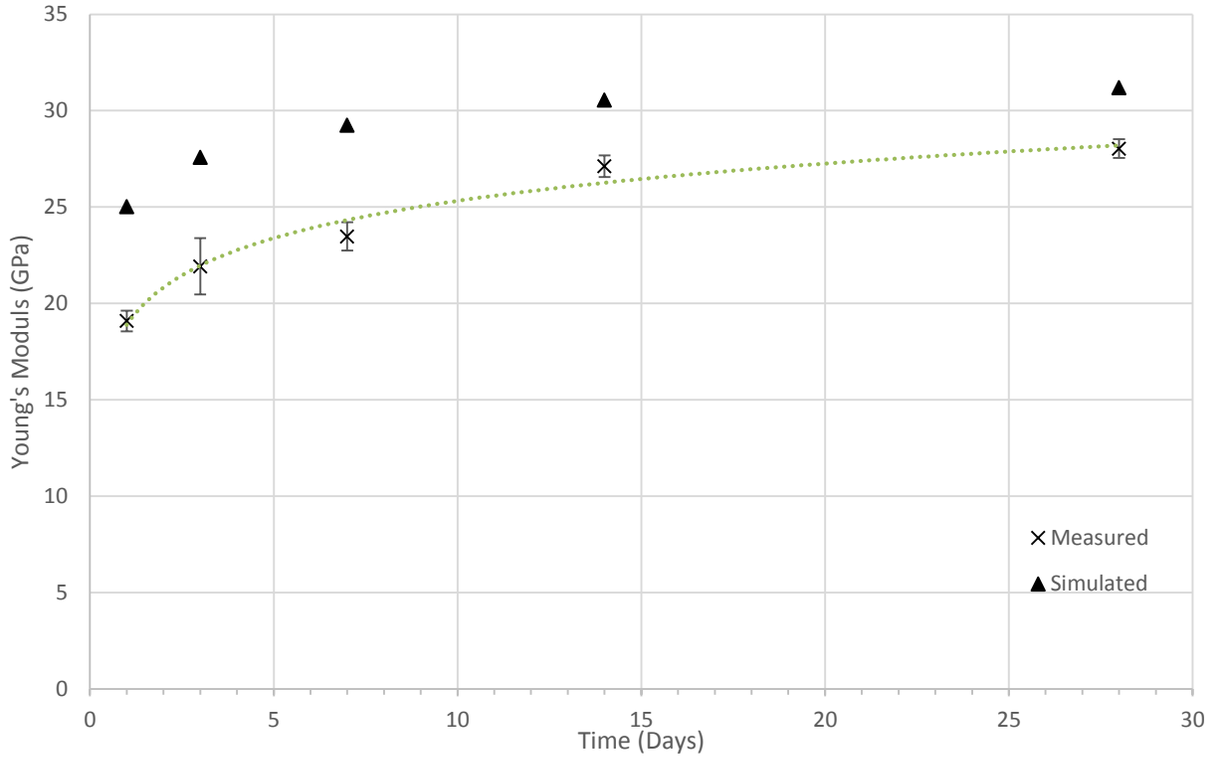


Figure 7-9: Young's Modulus vs. Time, w/c of 0.5

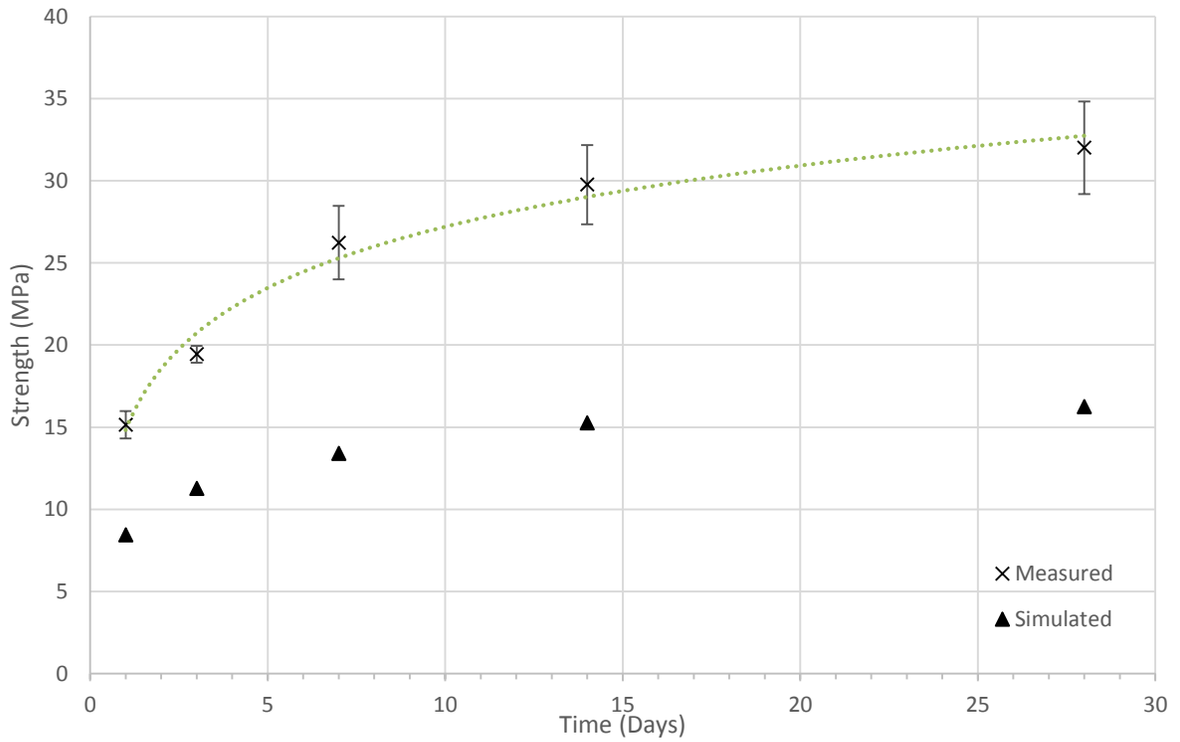


Figure 7-10: Compressive Strength vs. Time, w/c of 0.5

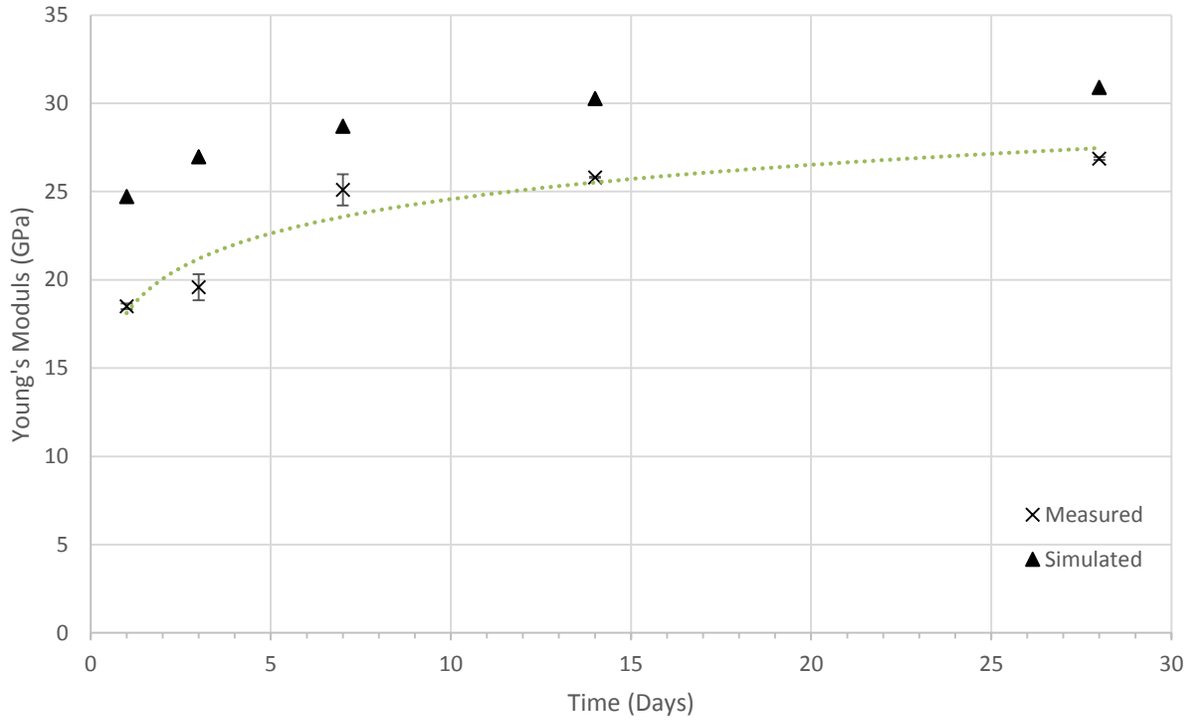


Figure 7-11: Young's Modulus vs. Time, w/c of 0.55

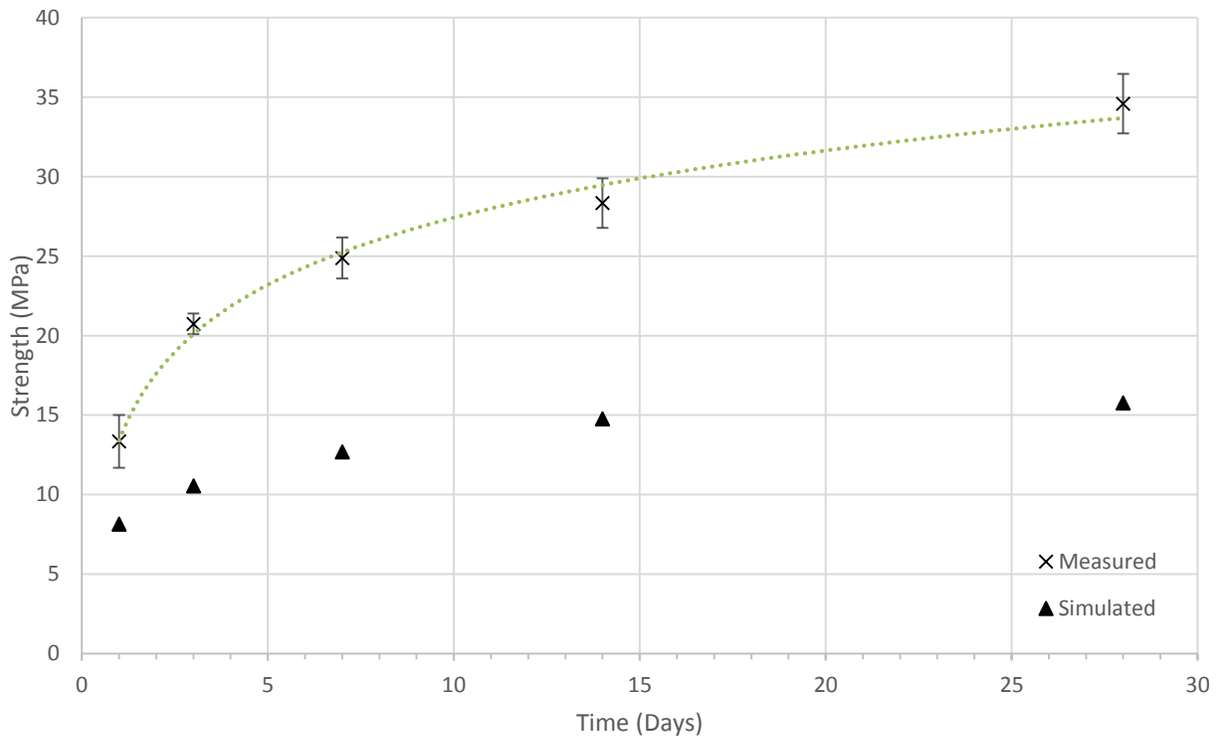


Figure 7-12: Compressive Strength vs. Time, w/c of 0.55

A comparison of Figures 7-1, 7-5, 7-7, and 7-9 indicates that the difference between the simulated and measured values of Young's modulus increased as w/c increased. The simulated values overestimated early-age modulus, while predicting later-age modulus closely at lower w/c. Simulated compressive strength was dramatically lower than measured compressive strength at all w/c and ages.

Input Sensitivity Study

Figures 7-13, 7-14, and 7-15 present the effects of the removal of cement calorimetry data, cement particle shape data, and aggregate shape data, respectively, relative to both the simulated control sample with all available inputs and the measured values for the physical control sample.

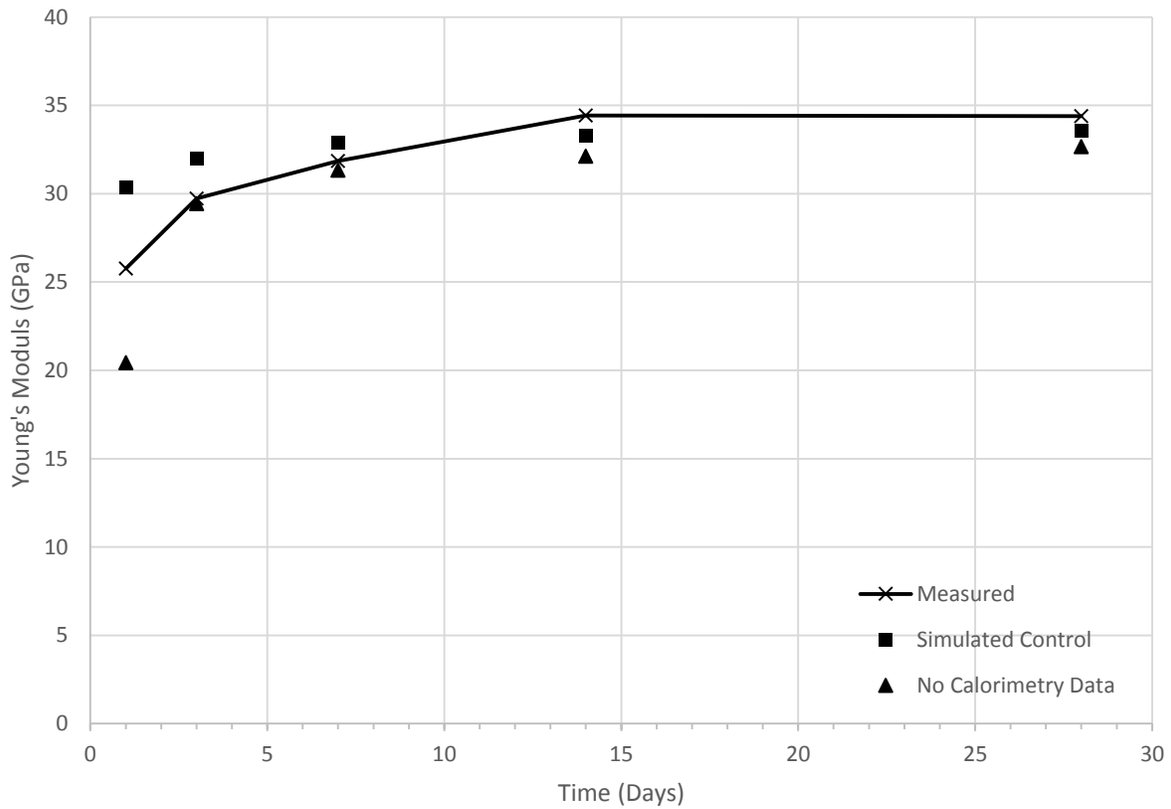


Figure 7-13: Young's Modulus vs. Time, Removal of Calorimetry Data

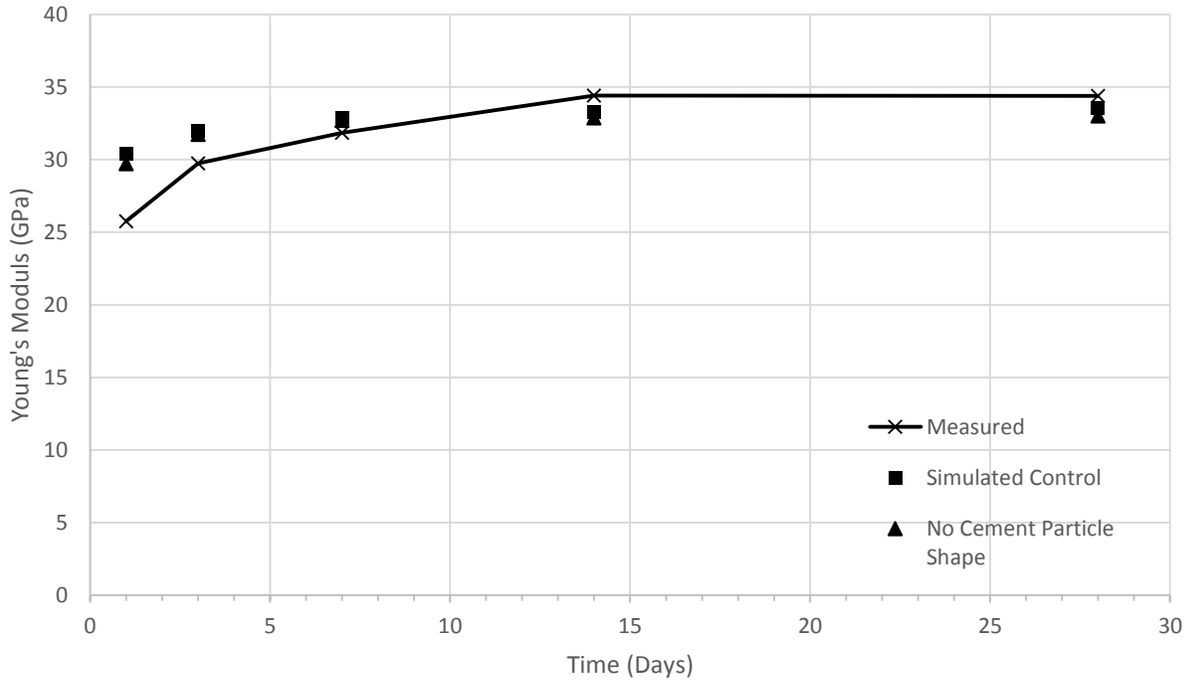


Figure 7-14: Young's Modulus vs. Time, Removal of Cement Particle Shape Data

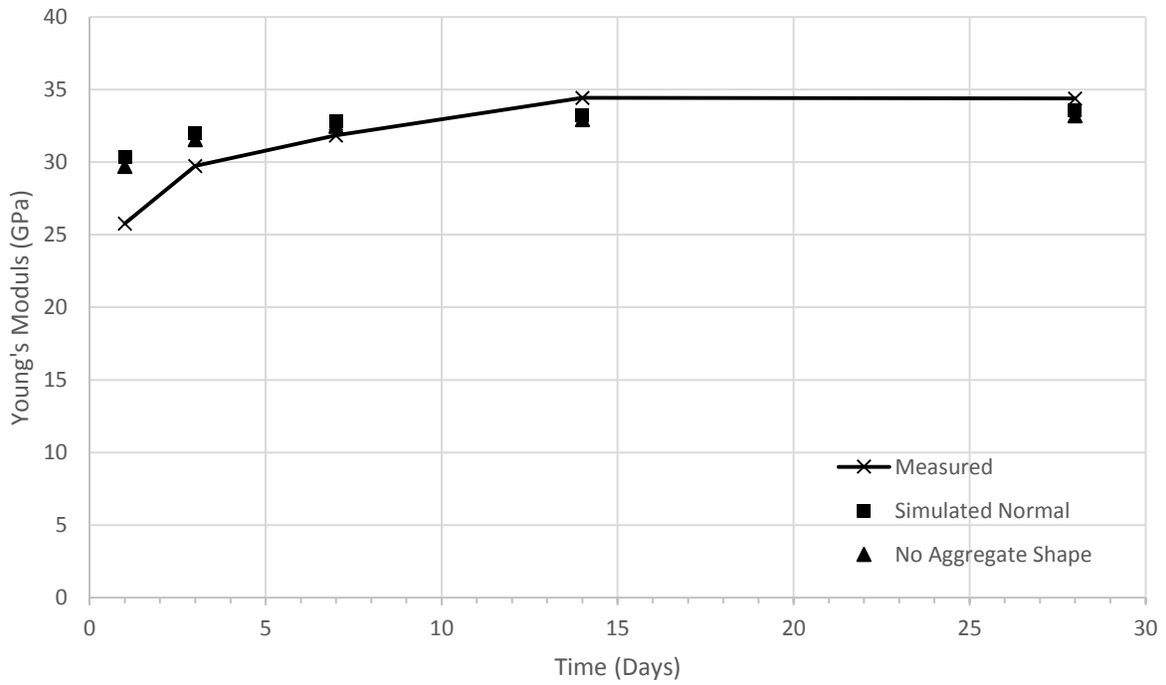


Figure 7-15: Young's Modulus vs. Time, Removal of Aggregate Shape Data

The absence of aggregate shape data and cement particle shape data (Figures 7-14 and 7-15) appeared to have very little influence on the simulated Young's modulus, while absence of calorimetry data (Figure 7-13) had a more dramatic effect on modulus. The simulation performed without utilizing measured calorimetry data underestimated the Young's modulus at early age, which was in opposition to the simulation with the calorimetry data, which overestimated modulus at early ages. At later ages, both simulations underestimated the modulus, with the no-calorimetry-data simulation doing so to a slightly greater extent.

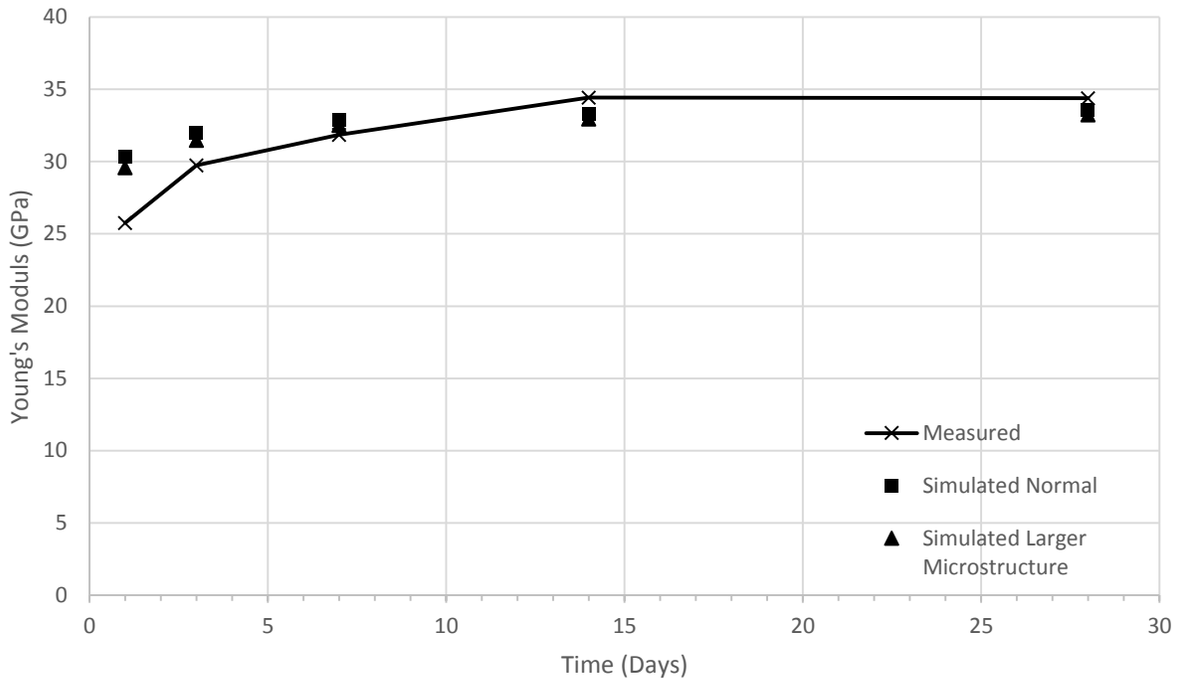


Figure 7-16: Young's Modulus vs. Time, Larger Virtual Microstructure

The effect on a simulation performed with a microstructure 50% larger on a side is shown by Figure 7-16. Similar to simulations performed without aggregate or cement particle shape, the larger microstructure had very little influence on the simulated results. Of note is the increase in the time required to run a simulation of a larger microstructure. The larger digital microstructure was 50% longer in each direction, resulting in a volume 3.375 times larger than the original. Thus the simulation of the larger digital microstructure took approximately 3.375 times longer.

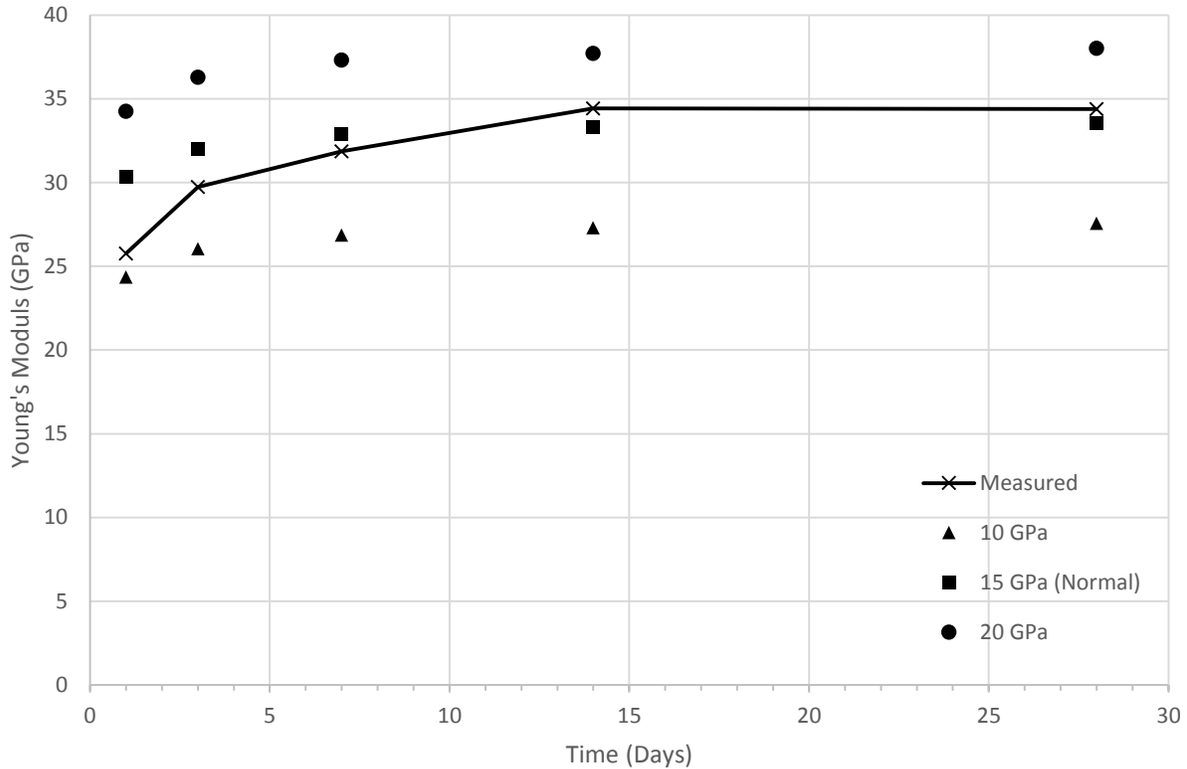


Figure 7-17: Young's Modulus vs. Time, Different Aggregate Moduli

The influence of aggregate moduli on modulus of elasticity of the simulated virtual microstructure is presented in Figure 7-17, which displays the results of simulations with three different aggregate moduli and the measured values. The influence of the elastic modulus of the aggregate on the simulated microstructure produced reasonable results. The simulated concrete modulus was directly proportional to the modulus used for the coarse aggregate. In other words, the use of a higher coarse aggregate modulus resulted in a higher simulated concrete modulus.

Three optional model input parameters, microstructure size, isothermal calorimetry, and cement particle shape, were found to be capable of influencing the formation or hydration of the virtual microstructure. To compare these parameters and their influence on the modeled results, simulations were performed to see their effects on the fraction solids connected, as presented in Figure 7-18.

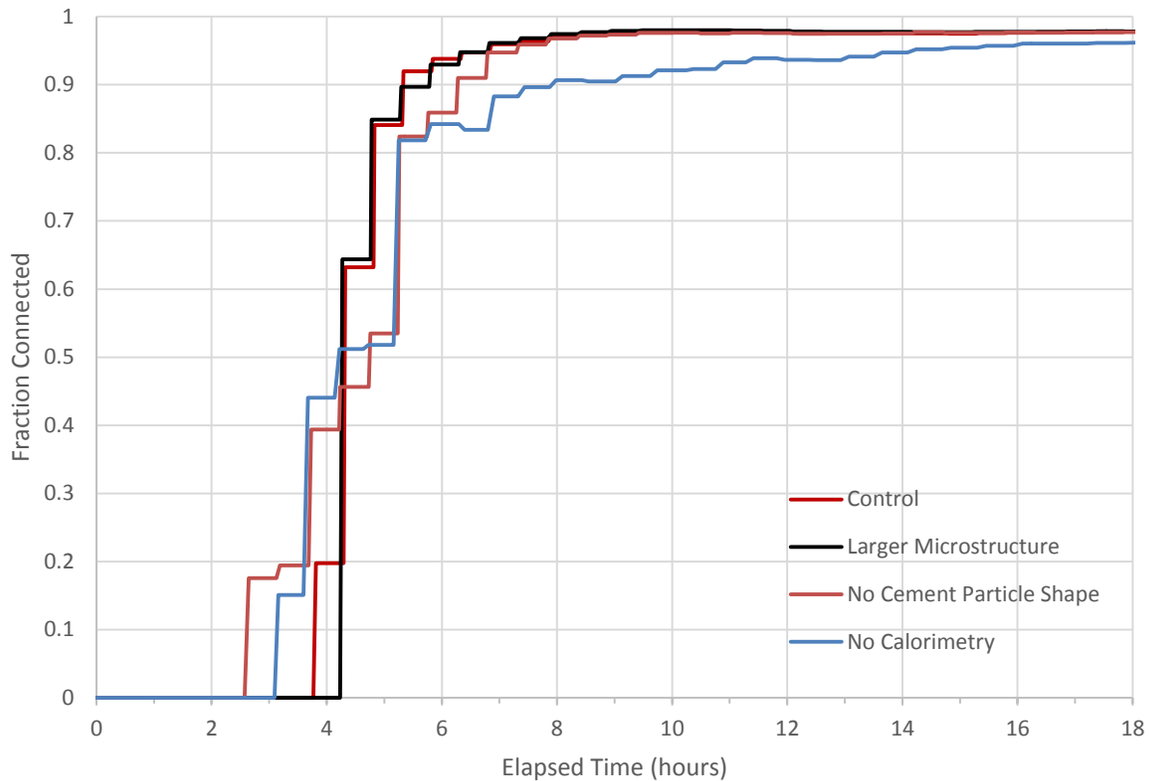


Figure 7-18: Fraction Solids Connected vs. Elapsed Time

Initial and final times of set are defined by their fraction of solids connected, 0.2 and 0.8, respectively. The results in Figure 7-18 indicate that the control simulation and the simulation with larger microstructure produce about the same fraction of solids connected with respect to time. Simulations performed in which the particle shape was not considered (default setting) produced a microstructure that resulted in an earlier initial time of set and delayed final time of set. The simulation performed with no isothermal calorimetry data (default setting) behaved in a similar manner, but to a lesser extent. The results indicate that simulations performed with default settings will not provide an accurate rate of development of the microstructure.

Admixture Time of Set

For the initial attempt to incorporate the effects of admixtures into VCCTL simulations, two assumptions were made: 1) the admixtures do not have a significant influence on the long-term cured microstructure, which determines properties such as strength and elastic modulus, and 2) the short-term effects can be accounted for by using the heat evolution-time

profile (from isothermal calorimetry) of the cementitious material plus admixtures instead of the profile for just the cementitious material. The typical reason for incorporating chemical admixtures into a concrete mixture is that the needed workability can be achieved with a reduced w/c. A reduced w/c is desirable because it results in increased strength due to reduced residual porosity.

The measured values for strength and elastic modulus for the mixes with different admixture additions are shown in Figures 7-19 and 7-20 below. Error bars for the no-admixture mix represent plus or minus one standard deviation of the sample data. Figures 7-19 and 7-20 indicate that the mix with 4 oz/cwt Type F admixture had a consistently higher strength and modulus at all ages. The other mixes attained similar strength to the control mix. It should be noted that even though the w/c was held constant at 0.4, the strength still significantly improved for the mix with 4 oz/cwt of Type F high-range water reducer. The improved strength and modulus indicates that the use of this water-reducing admixture improved the cured microstructure without a reduction in water (same w/c). It is apparent that the first assumption above (no significant change in the cured microstructure) may not always be valid.

Water-reducing admixtures improve the overall dispersion of particles in the mix, allowing for more efficient hydration and more uniform distribution of reaction products. This effect is accounted for in the VCCTL software by using the “dispersion” input parameter. As previously explained, the functional effect of enabling the dispersion input is that the assigned space between particles is more uniform, resulting in a more accurate simulation of the improved hydration that occurs due to the presence of a water-reducing admixture.

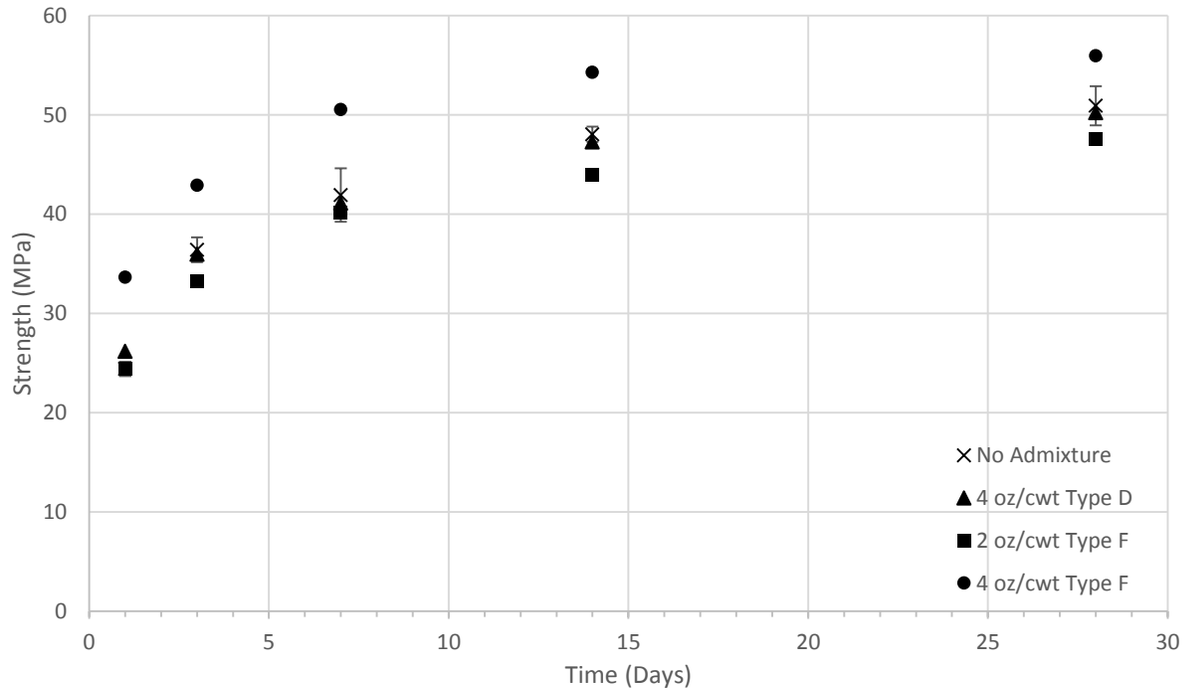


Figure 7-19: Measured Strength vs. Time for Different Admixtures, w/c = 0.4

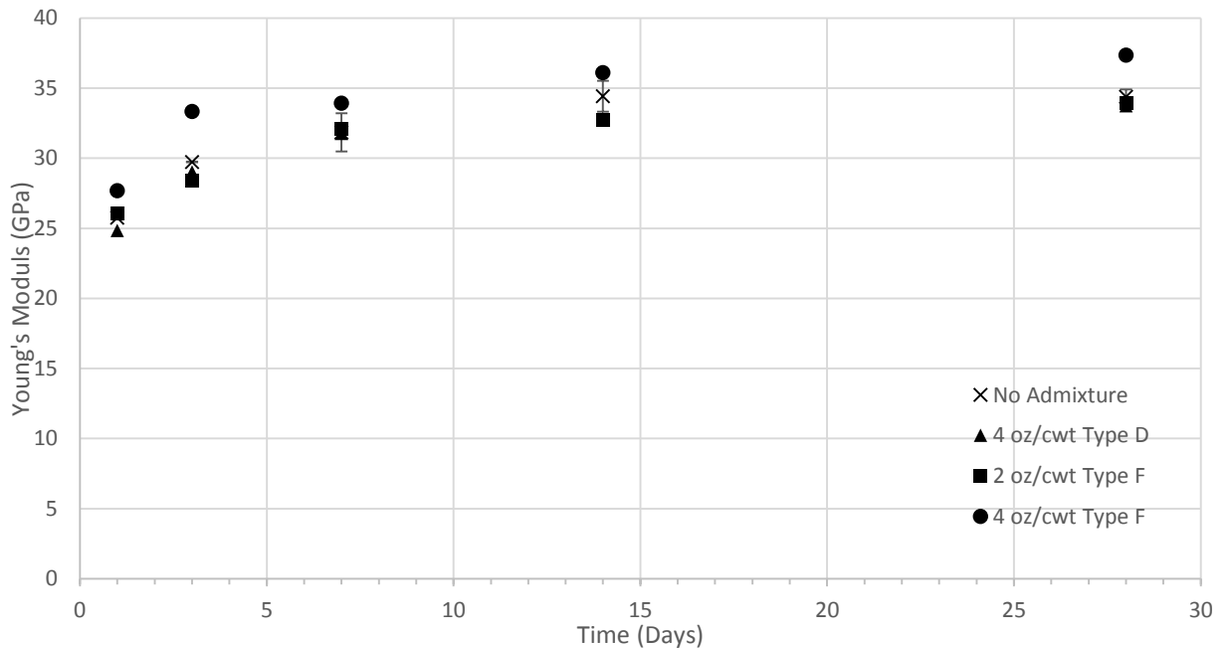


Figure 7-20: Measured Young's Modulus vs. Time for Different Admixtures, w/c = 0.4

Figures 7-21 through 7-26 present the results for strength and Young's modulus for simulated and measured mixes containing admixtures. For the comparison with the control mix, refer to Figures 7-5 and 7-6.

Figures 7-21 (4 oz/cwt Type D) and 7-22 (2 oz/cwt Type F) indicate that the model predicted the modulus at later ages to within one the standard deviation of measurements taken at the respective age. The model slightly underestimated the modulus for the mix with 4 oz/cwt of Type F (Figure 7-23). This corresponded with the increase in modulus as compared to the control mix in Figure 7-20. The compressive strength data shown in Figures 7-24 – 7-26 clearly indicate that the simulated compressive strength was considerably less than that of the measured compressive strength. As discussed later and shown in Figure 7-32, this large underestimation is mainly due to the particular equation used by the VCCTL software to calculate compressive strength.

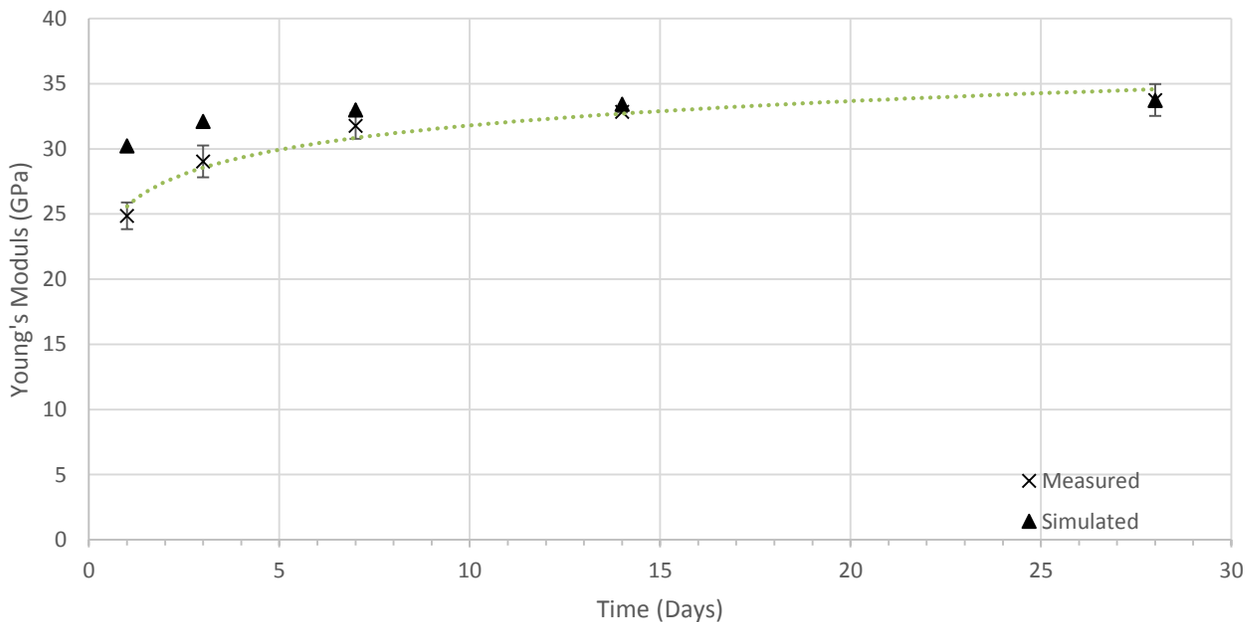


Figure 7-21: Young's Modulus vs. Time, 4 oz/cwt Type D Admixture, w/c = 0.4

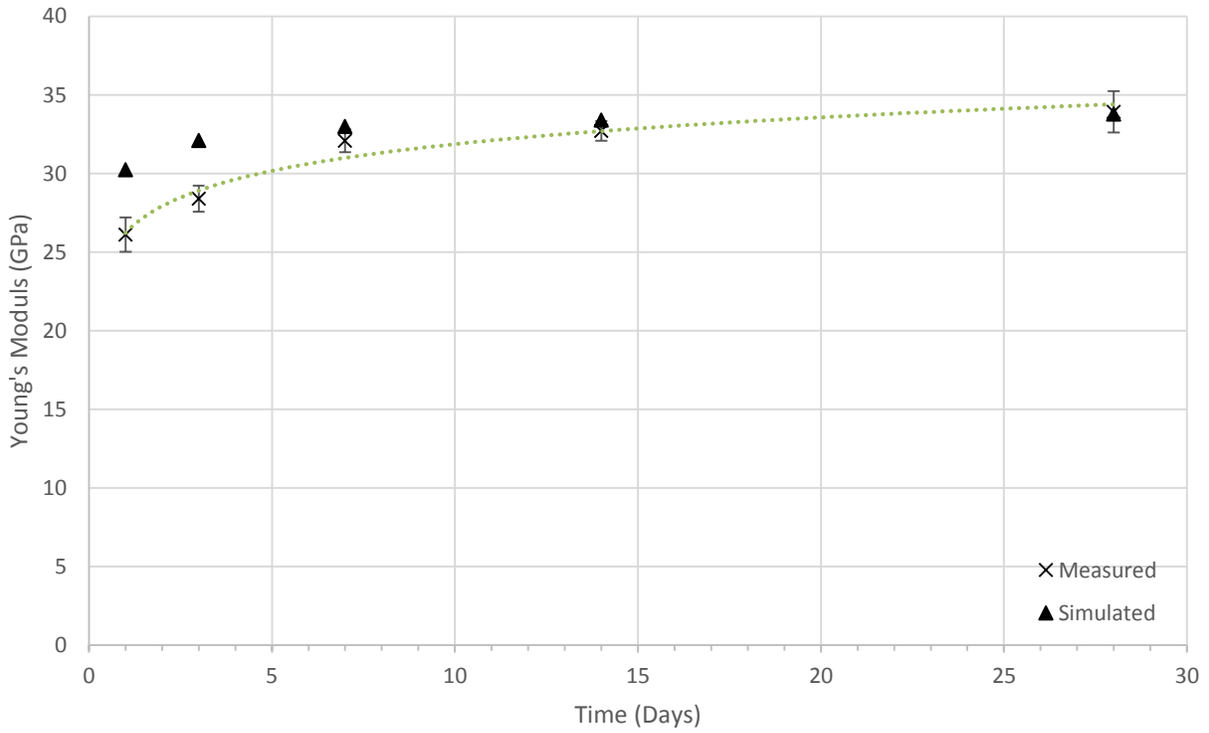


Figure 7-22: Young's Modulus vs. Time, 2 oz/cwt Type F Admixture, w/c = 0.4

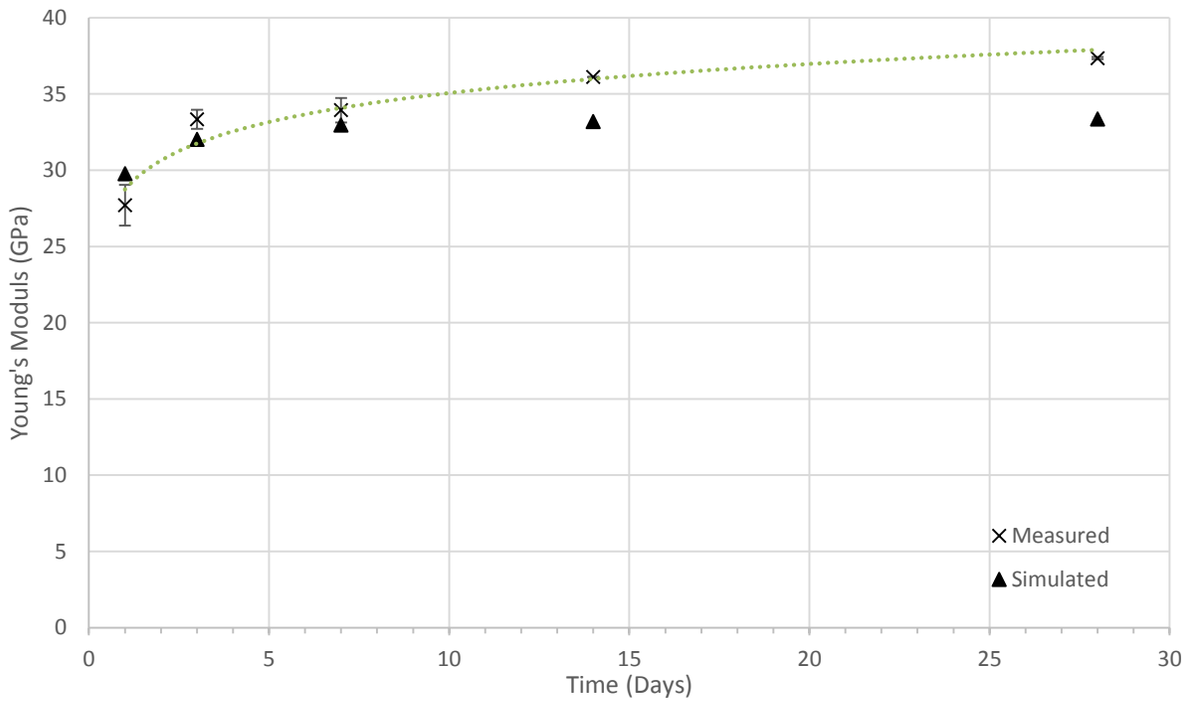


Figure 7-23: Young's Modulus vs. Time, 4 oz/cwt Type F Admixture, w/c = 0.4

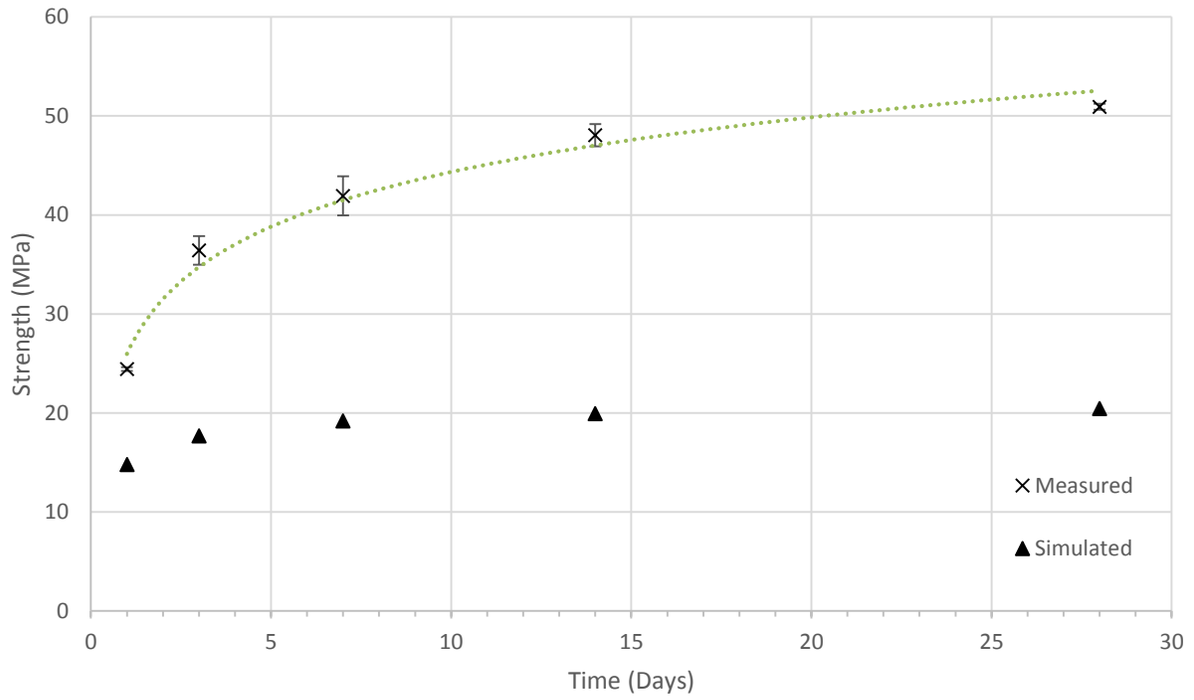


Figure 7-24: Strength vs. Time, 4 oz/cwt Type D Admixture, w/c = 0.4

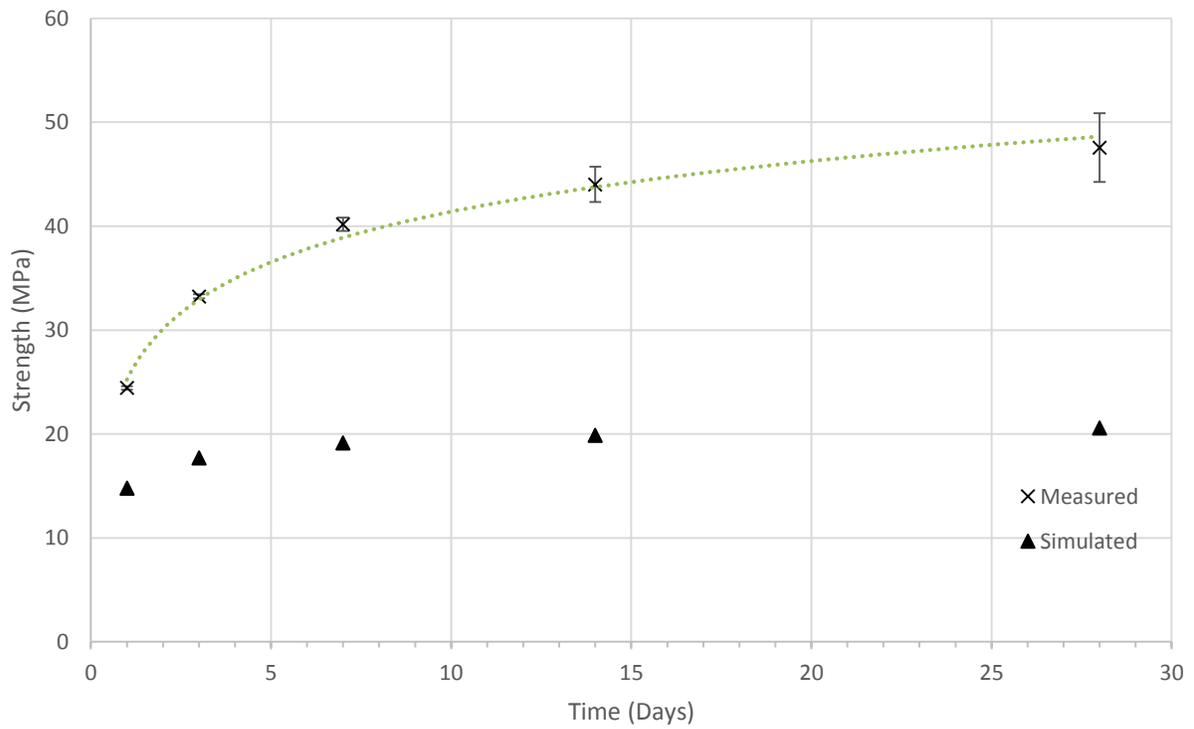


Figure 7-25: Strength vs. Time, 2 oz/cwt Type F Admixture, w/c = 0.4

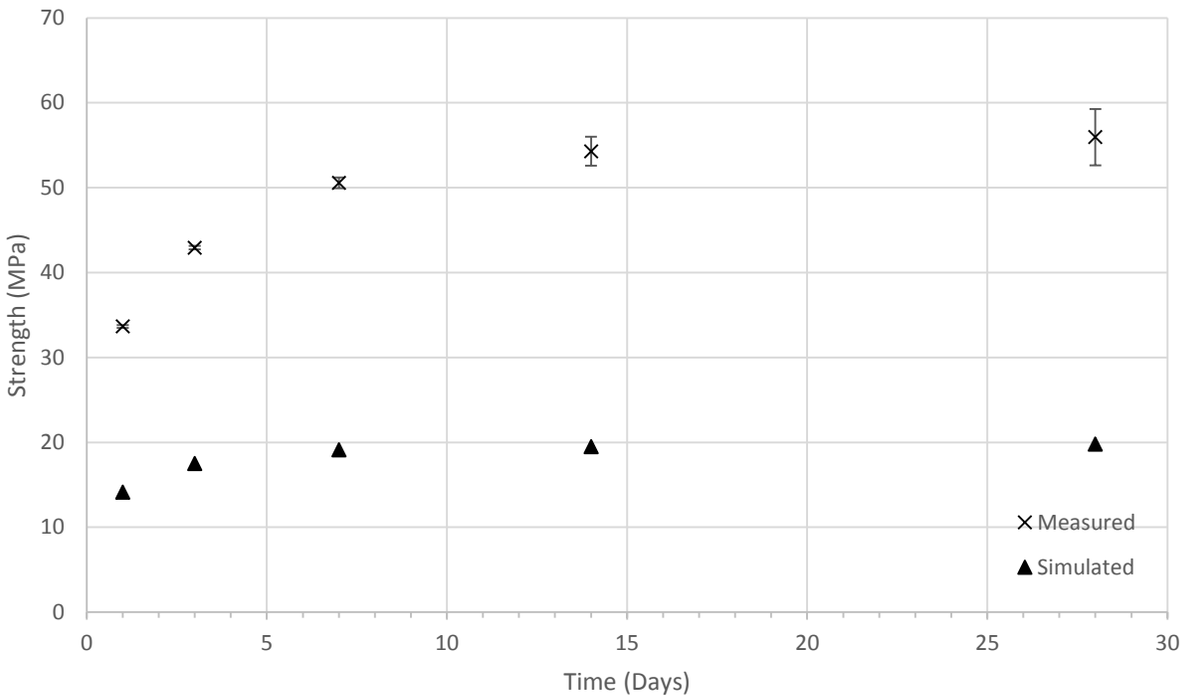


Figure 7-26: Strength vs. Time, 4 oz/cwt Type F Admixture, w/c = 0.4

The power and heat calorimetry curves used as inputs to simulate the mixes with admixtures are shown in Figures 7-27 and 7-28. The heat and power curves show different degrees of delay for the second peak that marks the start of the hydration of C_3S (Mindess et al, 2003) (at a time of approximately 9 hours). The power curve for the control mix produced a maximum value for the second peak approximately 10 hours after the introduction of the water, and the other mixes with 4 oz/cwt Type D and 2 oz/cwt Type F, and 4 oz/cwt of Type F experienced respective consecutive delays of approximately one hour each. The delay in power / heat generation is attributed to the repulsion of the cement grains at the time of admixture interaction, causing a delay in hydration (Mamlouk & Zaniewski, 2010).

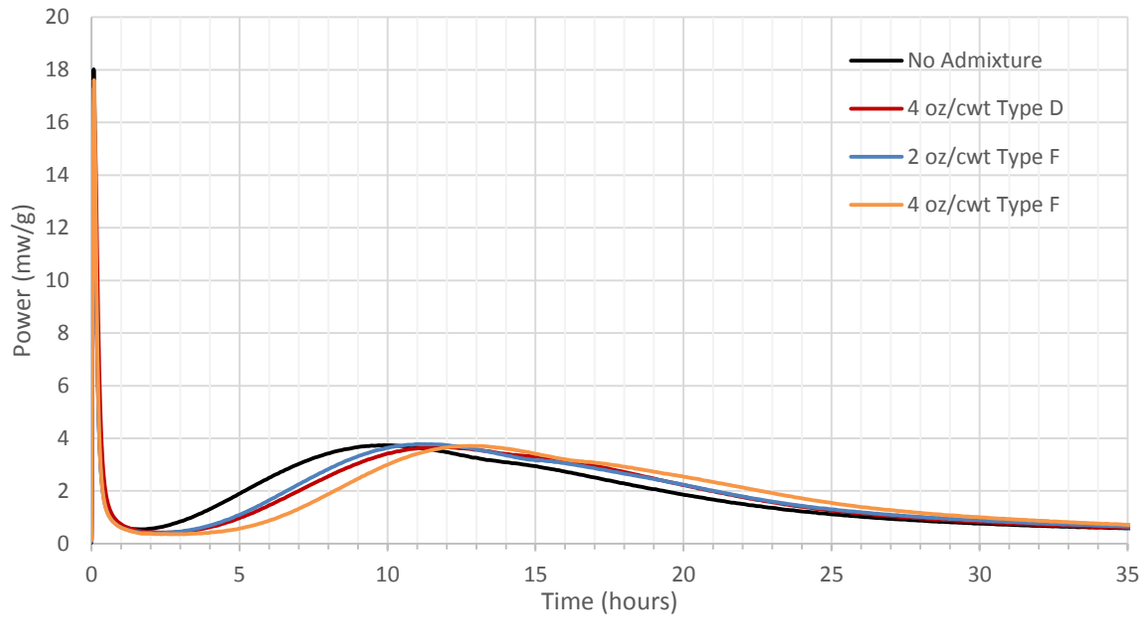


Figure 7-27: Power vs. Time

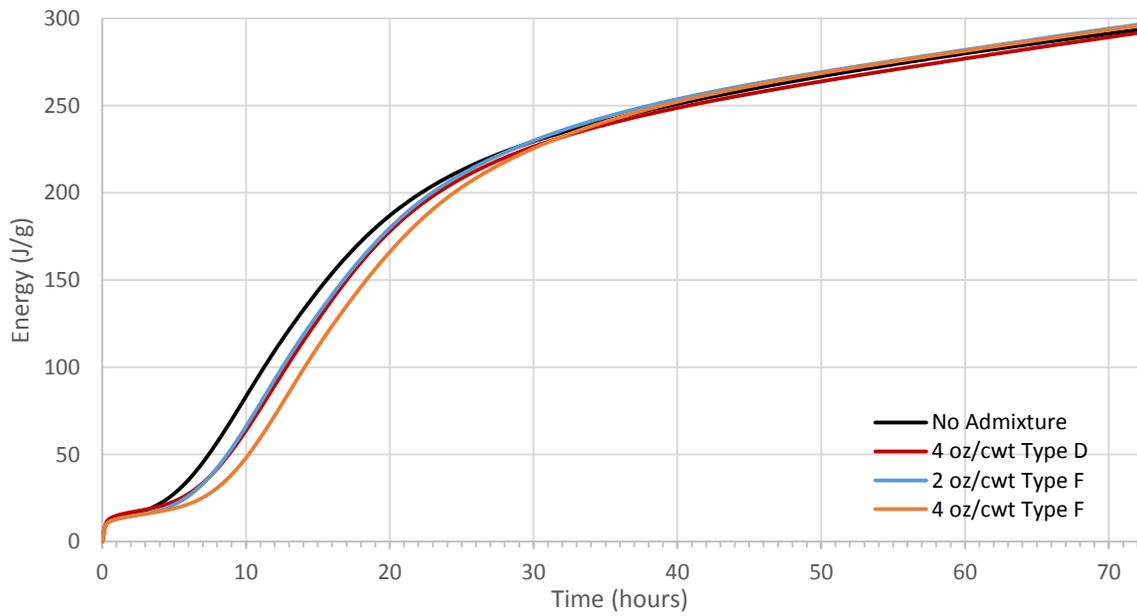


Figure 7-28: Energy vs. Time

The simulated fraction of solids connected for mixtures which incorporate admixtures is presented in Figure 7-29. The simulations incorporate the results of heat of hydration (isothermal calorimetry) results obtained in the laboratory as shown in Figures 7-27 and 7-28. The curves in Figure 7-29 provided by the simulation indicate that the rate of the connectivity of the solids

within the microstructure (fraction connected) is related to the rate and relative delay of hydration measured within the laboratory. This finding is of particular interest as it provided a potential technique for the incorporation of the liquid admixtures into simulations by the VCCTL.

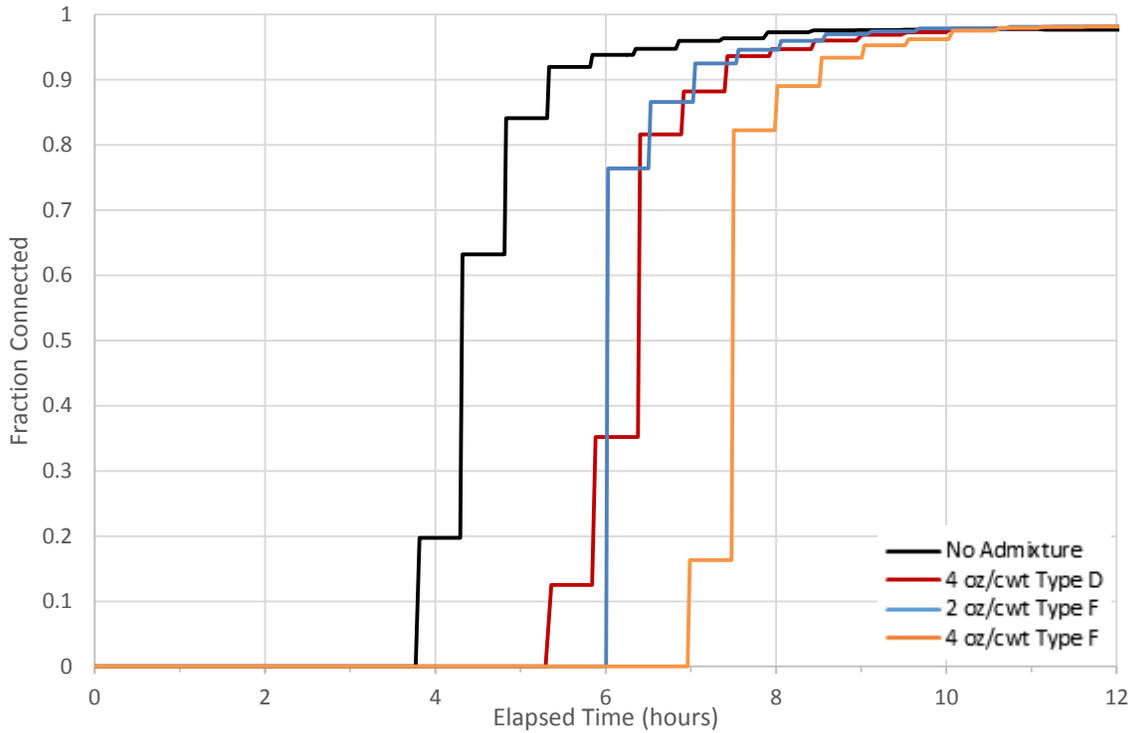


Figure 7-29: Fraction of Solids Connected vs. Time

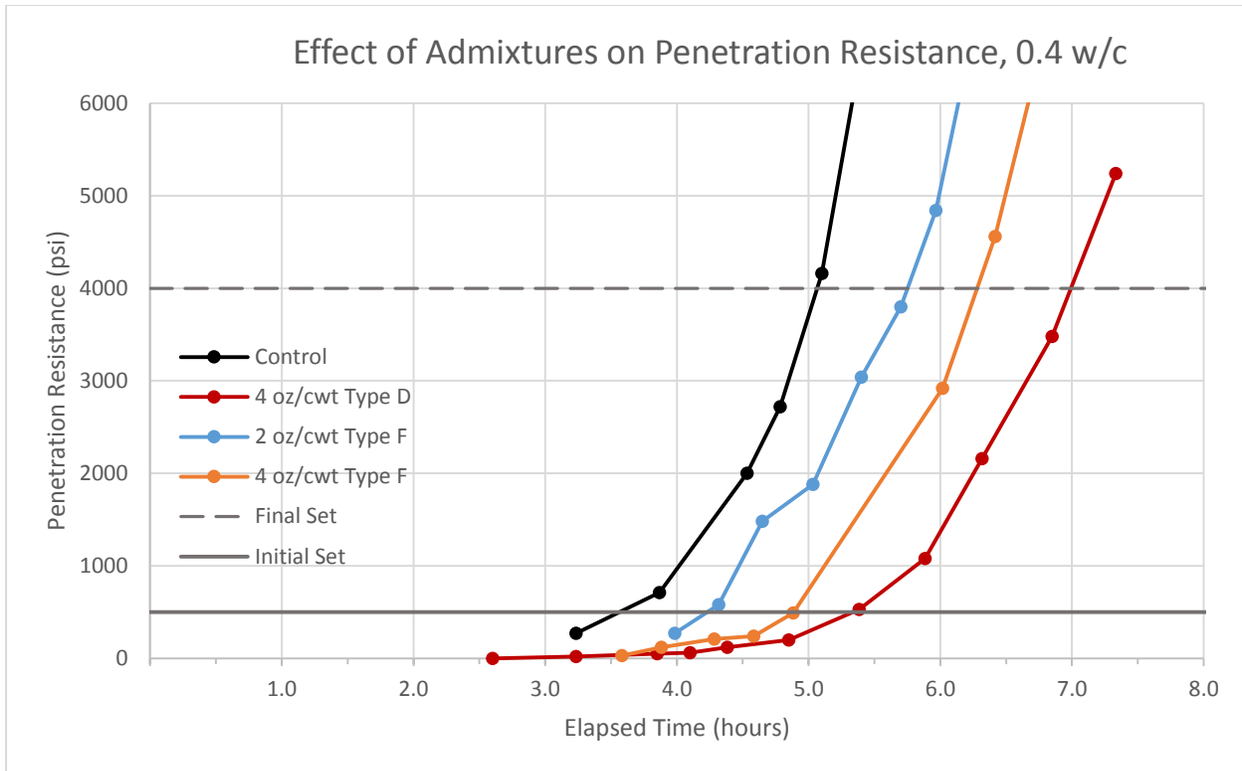


Figure 7-30: Penetration Resistance vs. Time, Different Admixtures and Dosages

The time of set via penetration resistance testing for the different mixes is presented in Figure 7-30. The results of the time of set testing indicate that the control mix and mixes with Type F admixture are in agreement with the results obtained in the isothermal calorimetry testing, with the exception of the Type D admixture (which incurred a longer delay of set time). The connectivity of solids, as obtained from the simulation, experienced a delay relative to the time of set as well. Thus, isothermal calorimetry and time of set testing provide potential input parameters for VCCTL for concrete mixtures which incorporate liquid admixtures. More research is needed to confirm this theory.

Discussion of Results

Modulus of Elasticity and Compressive Strength

The results from the w/c study mixes and simulations indicate that the VCCTL software makes reasonable predictions of modulus at a w/c of 0.4, with progressively more overestimation of modulus as the w/c increases.

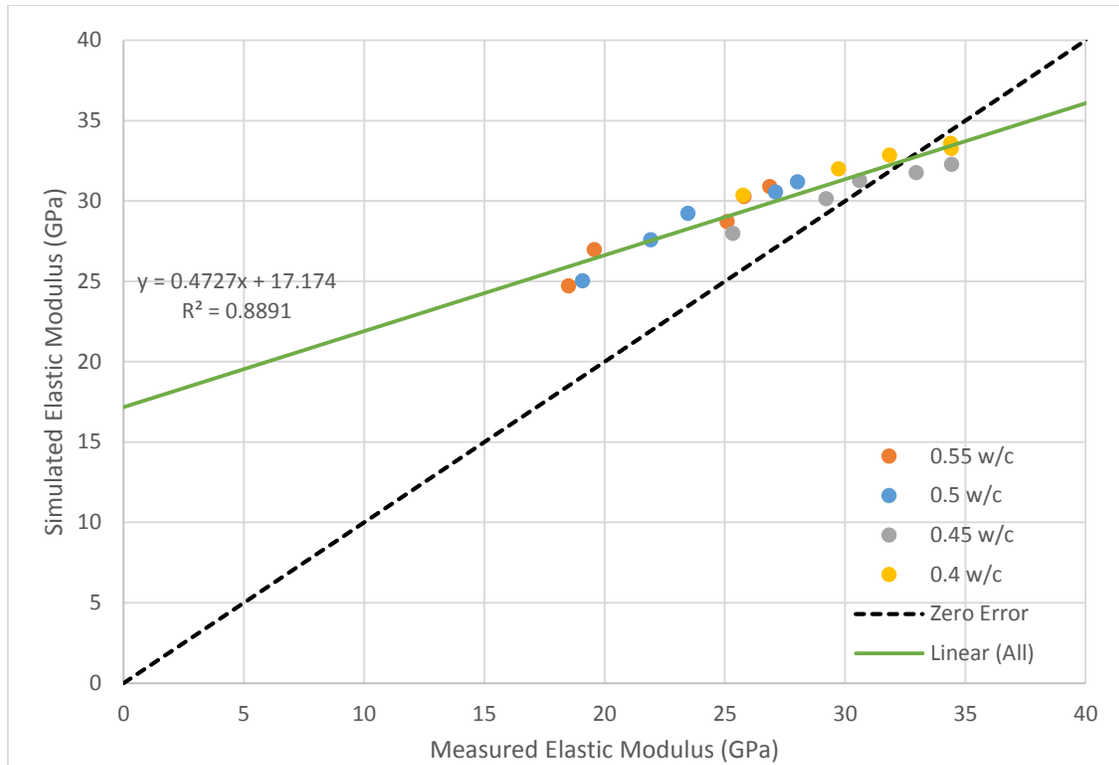


Figure 7-31: Simulated vs. Measured Elastic Modulus

The data presented in Figure 7-31 show the measured modulus of elasticity versus the simulated modulus of elasticity obtained for four different concrete mixtures at ages of 1, 3, 7, 14, and 28 days. The dashed line is a “zero error line” representing the ideal simulated values of elastic modulus with respect to experimentally measured values. The results indicated a general linear trend. The trendline for the data had an R^2 value of 0.89, with an intercept at 17.17 GPa, which corresponds to a significant overestimation of simulated modulus of elasticity versus measured modulus at low modulus values. The concrete samples which exhibited the lowest modulus of elasticity (measured) were early-age specimens that had a relatively high w/c. The modulus of elasticity is simulated by the VCCTL from the modulus of the virtual microstructure, and modulus development is directly related to the rate at which the virtual microstructure is hydrated. The data shown above indicate that the equation used by the VCCTL software to calculate the modulus for the virtual microstructure could be modified, using the linear relationship shown in Figure 7-31, to give modulus values that match those measured.

As discussed earlier, the heat of hydration data, as obtained via isothermal calorimetry, was used as a VCCTL input to provide a timeline to synchronize the rate of simulated microstructural development and corresponding heat release with measured values.

Regardless of the w/c or whether measured heat of hydration data was used for model calibration, the simulations of early-age modulus did not produce the level of accuracy as compared to later ages. Simulated moduli of elasticity at later ages were more comparable to measured values, with 28-day simulated values within five GPa of the measured values for each of the concrete mixtures. The exact cause of early-age discrepancies is difficult to determine directly; more research into the problem is required to illuminate the underlying reasons for this deficiency. When the cause is determined, investigations into potential corrections to the model can be performed.

The most obvious discrepancies between simulated and measured values occurred for strength. Strength was dramatically underestimated by the VCCTL software in all cases. Simulated modulus values were calculated from a simple empirical relationship. Figure 7-32 shows the relationship between modulus and strength for both simulated and measured values.

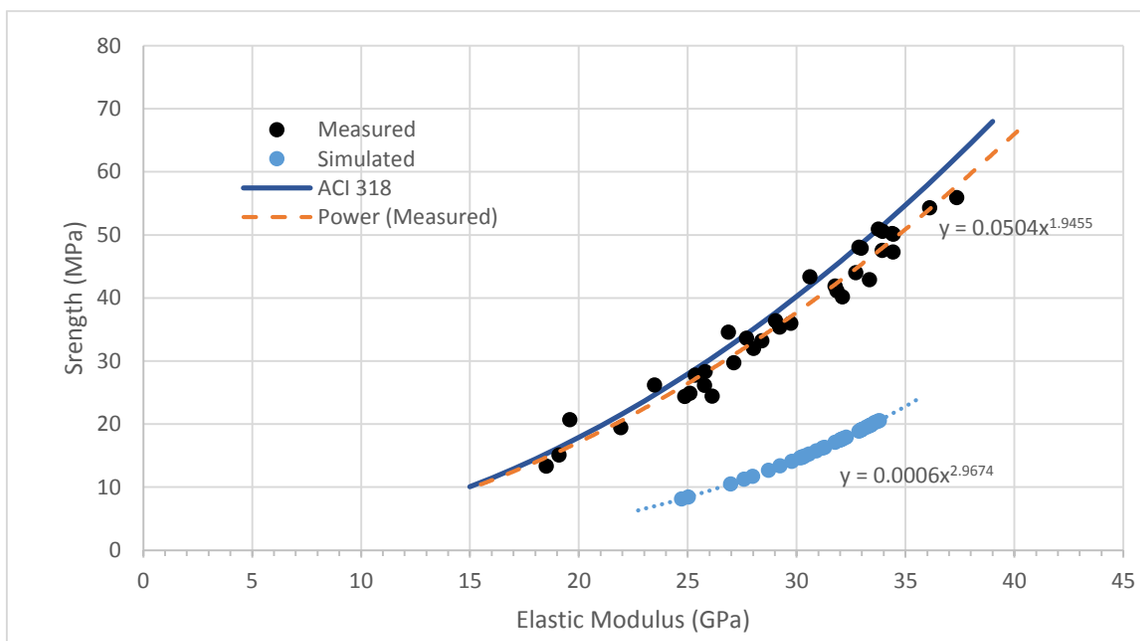


Figure 7-32: Strength vs. Elastic Modulus with VCCTL Software Power Fit and ACI 318

As previously discussed, Figures 7-24 – 7-26 provided results in which predicted compressive strengths obtained from the simulations were considerably underestimated with

respect to compressive strengths as obtained from the specimens tested in the laboratory. However, the discrepancy between simulated and measured compressive strength is due to the equation used calculate compressive strength from elastic modulus. The VCCTL uses empirical equations to calculate compressive strength from simulated modulus of elasticity. Equation 7-1 is the built-in equation for the VCCTL and originates from work performed by another member of the VCCTL consortium; however, no publications exist detailing the materials used in the concrete testing program from which this equation was developed. Equation 7-1 does not come close to fitting the actual data. Equation 7-2 provides the conversion from elastic modulus to compressive strength (in metric units) as per ACI 318 (Neville, 2011), and it can be seen that this equation fits the data fairly well. The best fit, calculated from the data, is given by Equation 7-3, and can be compared with the other two in Figure 7-32. Use of this equation in the VCCTL software for the conversion of compressive strength from elastic modulus would definitely improve the accuracy of the simulated values.

$$f'_c = 0.0006E_c^{2.9674} \quad (7-1)$$

$$f'_c = 0.0447E_c^2 \quad (7-2)$$

$$f'_c = 0.0504E_c^{1.9455} \quad (7-3)$$

Where:

E_c is modulus of elasticity in GPa and

f'_c is compressive strength in MPa

Though the relationship defined in ACI 318 appears as though it would predict strength fairly well at later ages in this circumstance, the use of empirical relationships does not take into account many of the factors that influence the behavior of concrete. It is possible that the discrepancy between the VCCTL software's predictions and the actual physical results is due to one of these factors, the most likely of which being the scenario in which the coarse aggregates being modeled do not have the same properties as those of the materials used to develop the empirical relationship.

Due to the lack of information on the materials used to develop the empirical relationship between modulus and strength, any possible explanation for the drastic underprediction of

strength for the concrete used in this study must come from specific knowledge about the properties of the materials used as well as an assumption about the nature of the materials used to establish the equation 7-1.

The concrete in this study exhibited fracture behavior that corresponds to the commonly described mechanism of failure for concrete. Typical concrete fails in compression when microcracks initiated at the interfacial transition zones surrounding aggregates propagate through the specimen, resulting in failure planes that pass around aggregate particles (Maso, 1996). This is called intergranular fracture. In the concrete used in this study however, the failure planes passed through the aggregate particles, commonly referred to as transgranular fracture. Figure 7-33 shows an example of a failed cylinder that exhibits this behavior. Coarse aggregate particles that have fractured are clearly visible in the two fragments as shown in Figure 7-33. An aggregate particle face in one fragment is mirrored in the other fragment, which came from the opposite side of the failure plane. Based on this phenomenon, it can be concluded that the strength of the interfacial transition zone in the concrete used in this research is sufficient to cause fracture to occur through the grain rather than through the interfacial transition zone.

Factors favoring transgranular fracture are better bonding between the aggregate and paste, weaker aggregate, or some combination of these two factors. The coarse aggregate used in this study was observably porous and angular, with an absorption of approximately 4.5%. The diffusion of hydration products into porous aggregates, which are initially saturated (all aggregate was in a saturated state) as well as the mechanical interlock between the cement paste and irregular aggregate surfaces can strengthen the interfacial transition zone to the point where it is no longer vulnerable to microcrack propagation (Maso, 1996). It has also been observed petrographically that the rough surfaces of Florida limestone (also referred to as limerock) aggregates provide good mechanical interlock and exhibit chemical bonding with cement paste (McClellan, et al., 1993).



Figure 7-33: Crushed Cylinder Fragments

The VCCTL software's underestimation of strength could be partially explained by the atypical bonding behavior of Florida limestone aggregate compared to the aggregate used by Siam Cement Group to develop the strength prediction formula. The modulus of concrete is measured within the linear elastic range, which is before microcracks begin to form and the condition of the aggregate interfacial transition zone (ITZ) becomes relevant (Neville, 2011). Thus it is possible to have two concretes that have the same modulus, even though the interfacial properties of the aggregates in those concretes are dramatically different. It follows that if given two concretes with the same elastic modulus, one with a typical aggregate interfacial zone and one with an interface similar to that which forms with Florida limestone, the concrete with typical aggregate interfacial behavior would develop microcracks more rapidly, resulting in earlier compressive failure. This could be a practical explanation for part of the VCCTL software's dramatic underestimation of strength, even while predicting elastic modulus to within one standard deviation of sample data values.

Model Sensitivity

The results of the investigation into the sensitivity of the model to the presence of different material input data show that several of the non-essential input values have little influence on the outcome of simulations. Aggregate shape data, while documented to have an influence on the strength of concrete (Jones & Kaplan, 1957), has no effect on the simulated strength because it has no effect on elastic modulus, the value from which strength is calculated. The size of the simulated microstructure had a very slight effect on the results of simulations in this case, though even a small increase in size results in a dramatic increase in simulation computing time.

The influence of aggregate elastic properties on the elastic properties of the concrete is substantial. The aggregate makes up a large fraction of the concrete by volume, and the VCCTL software calculates the modulus of the concrete based on the computed modulus of the paste as well as the relative volume fractions and elastic properties of the coarse and fine aggregate.

The absence of isothermal calorimetry data resulted in an underestimation of modulus, as opposed to an overestimation of modulus when calorimetry data was used as an input parameter for the simulated concrete. The absence of cement particle shape data had no appreciable influence on modulus, but had a similar effect on set time as the absence of calorimetry data. It is possible that if both of these inputs were missing the effect on set time could be compounded.

Admixture Influence on Set Time

The simulation of the delay in set time due to admixtures, by using isothermal calorimetry data obtained from those admixtures, was a partial success. The predicted delay in set time of the mixes containing two different dosage rates of Type F matched up with the measured set time delay of the actual concrete. The mix containing Type D did not however, with the predicted set occurring in roughly half the time of the actual measured set. Heat evolution and set time of concrete are two individual phenomena, and while they may be linked, the beginning of one does not necessarily imply the start of the other.

CHAPTER 8 CONCLUSIONS AND RECOMMENDATIONS

Conclusions

The results of this research led to the following conclusions about the VCCTL software:

1. The VCCTL software was found to be an effective tool for the simulation of elastic modulus of portland cement concrete, provided the materials being simulated are properly characterized.
2. At the current stage of development, the VCCTL software did not accurately predict the compressive strength of portland cement concrete using Florida limestone coarse aggregate. This is expected to be resolved by improving the accuracy of the raw material property inputs, and by making modifications to the software programming.
3. The VCCTL software currently does not have a means to incorporate the effects of admixtures on cement hydration. An initial attempt to integrate the effects of a water-reducing admixture, using heat of hydration data, was successful for Type F water reducer, but the software underestimated the setting time for Type D water reducer. Further research should be performed to reliably incorporate the effects of admixtures into the VCCTL.
4. The process of acquiring cement phase volume and surface area fraction data has been improved substantially through the use of automated scanning electron microscopy. This has resulted in a more efficient process to obtain cement characterization data for use in the VCCTL software.
5. The empirical predictions for compressive strength, which are based on elastic modulus and developed for concretes using coarse aggregates that were mineralogically and/or microstructurally different than typical Florida limestone aggregates, were not accurate for concretes made with Florida limestone. More research is needed to accurately predict the compressive strength based on the elastic properties of concrete containing Florida limestone coarse aggregates for different concrete mixtures.
6. Comparison of isothermal calorimetry data and corresponding time of set data has shown that a typical Type F high-range water-reducing admixture (superplasticizer) delayed time of set and shifted the main silicate hydration peak by the same amount of time. At the dosages explored within this study, the delay was proportional to the dosage rate.

7. Higher dosage rates of Type F high-range water-reducing admixture (superplasticizer) typically resulted in an increase in both elastic modulus and compressive strength, when compared to the control mixes.

Recommendations

Based on the findings from this study, the following recommendations were made:

1. The conclusions drawn from the work performed in this research discovered several potential topics for future research, both to improve the accuracy of the model as well as to explore different material inputs and outputs.
2. The empirical compressive strength model built into the VCCTL software, which is based on elastic modulus, does not accurately predict compressive strength for concrete mixtures which incorporate well for Florida limestone. Generalized empirical predictions for strength of concrete are inherently limited due to the variability of the materials used for production. A more fundamental approach to the simulation of concrete strength should be investigated.
3. Detailed characterization of the elastic properties of Florida limestones used to produce coarse aggregates for portland cement concrete should be performed. A database of properties of concrete mix designs containing Florida aggregate for use with the VCCTL software and other projects should be created.
4. The VCCTL software supports the modeling of admixtures if the specific phase surface deactivation behavior of the admixture is known. Methods of obtaining this information, either through material testing, or possibly from the admixture manufacturer, should be investigated.
5. There are a number of materials that can be modeled in the VCCTL software that were not considered for this research. There is support for both fly ash and blast furnace slag hydration in the VCCTL software, though the accuracy of the model in this respect is largely unknown. The techniques required to characterize these materials are also more involved due to the significant glassy (amorphous) phase contents of their compositions. The methods by which these materials can be characterized and the accuracy with which they are simulated in the VCCTL software should be explored.

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APPENDIX A
 MATHEMATICAL EQUATIONS REFERENCE FOR THE CALCULATION OF ELASTIC
 MODULI USING DIFFERENTIAL EFFECTIVE MEDIUM THEORY

The equations used to create a single effective particle are as follows

$$2A(G/G_2) + 2B(G/G_2) + C = 0$$

Where:

$$A = 8z(4 - 5\nu_2)\eta_\alpha p^{\frac{10}{3}} - 2[63z\eta_\beta + 2\eta_\alpha\eta_\beta]p^{\frac{7}{3}} + 252z\eta_\beta p^{\frac{5}{3}} - 50z(7 - 12\nu_2 + 8\nu_2^2)\eta_\beta p^{5/3} \\ + 4(7 - 10\nu_2)\eta_\gamma\alpha\eta_\beta$$

$$B = -2z(1 - 5\nu_2)\eta_\alpha p^{\frac{10}{3}} - 2[63z\eta_\beta + 2\eta_\alpha\eta_\gamma]p^{\frac{7}{3}} - 252z\eta_\beta p^{\frac{5}{3}} + 75z(3 - \nu_2)\eta_\beta\nu_2 p \\ + 3/2(15\nu_2 - 7)\eta_\gamma\alpha\eta_\beta$$

$$C = 4z(5\nu_2 - 7)\eta_\alpha p^{\frac{10}{3}} - 2[63z\eta_\beta + 2\eta_\alpha\eta_\gamma]p^{\frac{7}{3}} + 252z\eta_\beta p^{\frac{5}{3}} + 25z(\nu_2^2 - 7)\eta_\beta p + (7 \\ - 5\nu_2)\eta_\alpha\beta\eta_\beta$$

G = The effective shear modulus

K = The effective bulk modulus, given by:

$$K = K_2 + \frac{c(K_3 - K_2)}{1 + (1 - c)\frac{K_3 - K_2}{K_2 + \frac{4}{3}G_2}}$$

The differential equations used to solve numerically for the composite modulus are as follows

$$\frac{dK}{d\phi} = -kK/\phi$$

$$\frac{dG}{d\phi} = -gG/\phi$$

Where

$$k = \frac{(K_m + \frac{4}{3}G_m)(K_p + K_m)}{K_m(K_p + \frac{4}{3}G_m)}$$

$$g = \frac{(K_m + \frac{4}{3}G_m)(G_p + G_m)}{3G_m(K_m + \frac{8}{9}G_m) + 2G_p(K_m + 2G_m)}$$

m = Specified property of the matrix, and

p = Specified property of the particle.

APPENDIX B
SEM FIELD RAW DATA

Table B-1 Fields Acquired by the RJ Lee Group

Field	C ₃ S Area	Perimeter	C ₂ S Area	Perimeter	C ₃ A Area	Perimeter	C ₄ AF Area	Perimeter
1	0.58	0.31	0.23	0.51	0.10	0.15	0.08	0.04
2	0.58	0.31	0.23	0.50	0.10	0.14	0.08	0.04
3	0.50	0.33	0.26	0.41	0.07	0.15	0.18	0.11
4	0.60	0.36	0.23	0.46	0.01	0.01	0.16	0.17
5	0.43	0.31	0.36	0.50	0.08	0.09	0.14	0.10
6	0.47	0.22	0.33	0.52	0.13	0.19	0.07	0.07
7	0.41	0.25	0.29	0.39	0.16	0.29	0.15	0.06
8	0.51	0.36	0.27	0.37	0.08	0.12	0.14	0.15
9	0.46	0.33	0.33	0.45	0.06	0.12	0.15	0.11
10	0.52	0.50	0.29	0.31	0.10	0.13	0.08	0.06
11	0.58	0.32	0.22	0.45	0.05	0.05	0.15	0.18
12	0.45	0.29	0.31	0.41	0.09	0.20	0.15	0.09
Avg.	0.51	0.32	0.28	0.44	0.09	0.14	0.13	0.10

Table B-2: Fields Acquired by the University of Florida

Field	C ₃ S Area	Perimeter	C ₂ S Area	Perimeter	C ₃ A Area	Perimeter	C ₄ AF Area	Perimeter
1	0.47	0.26	0.32	0.66	0.05	0.06	0.13	0.01
3	0.47	0.22	0.45	0.71	0.01	0.02	0.07	0.04
5	0.41	0.16	0.34	0.64	0.10	0.11	0.17	0.08
6	0.57	0.34	0.27	0.38	0.11	0.19	0.03	0.06
9	0.52	0.51	0.28	0.31	0.02	0.01	0.18	0.16
Avg.	0.49	0.31	0.32	0.54	0.06	0.07	0.11	0.07